Experimental Cross Sections for Reactions of Heavy Ions and ²⁰⁸Pb, ²⁰⁹Bi, ²³⁸U, and ²⁴⁸Cm Targets

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Joshua Barnes Patin

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Abstract

Experimental Cross Sections for Reactions of Heavy Ions and ²⁰⁸Pb, ²⁰⁹Bi, ²³⁸U, and ²⁴⁸Cm Targets

by

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The study of the reactions between heavy ions and ²⁰⁸Pb, ²⁰⁹Bi, ²³⁸U, and ²⁴⁸Cm targets was performed to look at the differences between the cross sections of hot and cold fusion reactions. Experimental cross sections were compared with predictions from statistical computer codes to evaluate the effectiveness of the computer code in predicting production cross sections.

Hot fusion reactions were studied with the MG system, catcher foil techniques and the Berkeley Gas-filled Separator (BGS). 3n- and 4n-exit channel production cross sections were obtained for the ²³⁸U(¹⁸O,xn)^{256-x}Fm, ²³⁸U(²²Ne,xn)^{260-x}No, and ²⁴⁸Cm(¹⁵N,xn)^{263-x}Lr reactions and are similar to previous experimental results. The experimental cross sections were accurately modeled by the predictions of the HIVAP code using the Reisdorf and Schädel parameters and are consistent with the existing systematics of 4n exit channel reaction products.

Cold fusion reactions were examined using the BGS. The 208 Pb(48 Ca,xn) $^{256-x}$ No, 208 Pb(50 Ti,xn) $^{258-x}$ Rf, 208 Pb(51 V,xn) $^{259-x}$ Db, 209 Bi(50 Ti,xn) $^{259-x}$ Db, and 209 Bi(51 V,xn) $^{260-x}$ Sg

reactions were studied. The experimental production cross sections are in agreement with the results observed in previous experiments. It was necessary to slightly alter the Reisdorf and Schädel parameters for use in the HIVAP code in order to more accurately model the experimental data. The cold fusion experimental results are in agreement with current 1n- and 2n-exit channel systematics.

Dedication

This dissertation is dedicated to my mom and dad. Without your constant love, support, and guidance, none of this would have ever been possible. I love you both!

Table of Contents

1	Introd	duction				
	1.1	Discovery of the transuranium elements	1			
	1.2	Cold and hot fusion	14			
	1.3	Scope	26			
2	Evapo	oration Codes				
	2.1	JORPLE and SPIT	27			
	2.2	HIVAP	30			
3	Exper	imental procedures	34			
	3.1	MG system	36			
	3.2	Catcher foil experiments	42			
	3.3	Berkeley Gas-filled Separator experiments	48			
4	Exper	imental results	59			
	4.1	248 Cm + 15 N	61			
		4.1.1 ²⁶⁰ Lr	62			
		4.1.2 ²⁵⁹ Lr	72			
	4.2	$^{238}\text{U} + ^{18}\text{O} \rightarrow ^{252}\text{Fm} + 4\text{n}$	75			
	4.3	$^{238}U + ^{22}Ne$	87			
		4.3.1 ²⁵⁷ No	95			
		4.3.2 ²⁵⁶ No	97			
	4.4	208 Pb + 48 Ca	102			
		4.4.1 ²⁵⁵ No and ²⁵⁴ No	106			
		4.4.2 ²⁵³ No and ²⁵² No	116			

	4.5	$^{208}\text{Pb} + ^{50}\text{Ti}$	122
		4.5.1 ²⁵⁷ Rf	124
		4.5.2 ²⁵⁶ Rf	130
	4.6	$^{208}\text{Pb} + ^{51}\text{V} \rightarrow ^{257}\text{Db} + 2\text{n}$	135
	4.7	209 Bi + 50 Ti	139
		4.7.1 ²⁵⁸ Db and ²⁵⁷ Db	141
	4.8	$^{209}\text{Bi} + ^{51}\text{V} \rightarrow ^{258}\text{Sg} + 2\text{n}$	146
5	Discus	ssion	
	5.1	Cross sections	153
	5.2	Exit channel systematics	170
	5.3	HIVAP cross sections	175
	5.4	Odd-particle effects	182
6	Concl	usions and future research	
	6.1	Conclusions	187
	6.2	Future research	190
Apper	ndix A		192
Apper	ndix B		219
Refere	ences		220

List of Figures

1.1	Graphs of projectile (ion) mass versus E* _{min}	15
1.2	Graph of minimum excitation energy versus projectile mass	16
1.3	1n-exit channel cross section systematics	19
1.4	Graph of production cross sections versus atomic number	22
1.5	Graph of cold fusion cross sections versus atomic number	23
1.6	Cross sections versus element Z comparison	24
3.1	Target chamber for the MG experimental procedure	37
3.2	MG wheel collection and detection system	38
3.3	Catcher foil target chamber	44
3.4	Chemical separation flowchart	46
3.5	BGS schematic	49
3.6	Drawing of the Rutherford detectors	52
4.1	Sum spectrum from the 75.9 MeV ¹⁵ N on ²⁴⁸ Cm experiment	63
4.2	Sum spectrum from the 78.8 MeV ¹⁵ N on ²⁴⁸ Cm experiment	64
4.3	Cross sections from the ²⁴⁸ Cm(¹⁵ N,3n) ²⁶⁰ Lr reaction	71
4.4	Cross sections from the ²⁴⁸ Cm(¹⁵ N,4n) ²⁵⁹ Lr reaction	74
4.5	Sum spectrum for the 82.4 MeV experiment	78
4.6	Sum spectrum for the 94.9 MeV experiment	79
4.7	Sum spectrum for the 99.3 MeV experiment	80
4.8	Cross sections from the ²³⁸ U(¹⁸ O,4n) ²⁵² Fm reaction	86
4.9	214 Ac decay curve measurement, 118 MeV 22 Ne + 390 μ g/cm 2 238 U	92
4.10	²¹⁴ Ac growth curve measurement, 118 MeV ²² Ne + 390 μg/cm ^{2 238} U	93

4.11	²¹⁴ Ac growth curve measurement, 108 MeV ²² Ne + 100 μg/cm ^{2 238} U	94
4.12	Cross sections from the ²³⁸ U(²² Ne,3n) ²⁵⁷ No reaction	96
4.13	Cross sections from the ²³⁸ U(²² Ne,4n) ²⁵⁶ No reaction	101
4.14	EVR-α correlation spectra from the ²⁰⁸ Pb(⁴⁸ Ca,2n) ²⁵⁴ No reaction	108
4.15	Cross sections from the ²⁰⁸ Pb(⁴⁸ Ca,1n) ²⁵⁵ No reaction	114
4.16	Cross sections from the ²⁰⁸ Pb(⁴⁸ Ca,2n) ²⁵⁴ No reaction	115
4.17	Cross sections from the ²⁰⁸ Pb(⁴⁸ Ca,3n) ²⁵³ No reaction	120
4.18	Cross sections from the ²⁰⁸ Pb(⁴⁸ Ca,4n) ²⁵² No reaction	121
4.19	EVR-α correlation spectra from the ²⁰⁸ Pb(⁵⁰ Ti,1n) ²⁵⁷ Rf reaction	125
4.20	α - α correlation spectra from the 208 Pb(50 Ti,1n) 257 Rf reaction	127
4.21	Cross sections from the ²⁰⁸ Pb(⁵⁰ Ti,1n) ²⁵⁷ Rf reaction	129
4.22	EVR-SF correlation spectra from the ²⁰⁸ Pb(⁵⁰ Ti,2n) ²⁵⁶ Rf reaction	133
4.23	Cross sections from the ²⁰⁸ Pb(⁵⁰ Ti,2n) ²⁵⁶ Rf reaction	134
4.24	Cross sections from the ²⁰⁸ Pb(⁵¹ V,2n) ²⁵⁷ Db reaction	138
4.25	Cross sections from the ²⁰⁹ Bi(⁵⁰ Ti,1n) ²⁵⁸ Db reaction	144
4.26	Cross sections from the ²⁰⁹ Bi(⁵⁰ Ti,2n) ²⁵⁷ Db reaction	145
4.27	Cross sections from the ²⁰⁹ Bi(⁵¹ V,2n) ²⁵⁸ Sg reaction	151
5.1	Gaussian fits to the $^{208}\text{Pb} + ^{50}\text{Ti}$ and $^{209}\text{Bi} + ^{50}\text{Ti}$ reactions from GSI	157
5.2	Gaussian fits to the experimental $^{208}\text{Pb} + ^{50}\text{Ti}$ and $^{209}\text{Bi} + ^{50}\text{Ti}$ reactions	160
5.3	Gaussian fits to the experimental $^{208}\text{Pb} + ^{48}\text{Ca}$ and $^{209}\text{Bi} + ^{51}\text{V}$ reactions	161
5.4	Gaussian fit to the experimental ²⁰⁸ Pb + ⁵¹ V reaction	162
5.5	Gaussian fits to the experimental $^{238}\text{U} + ^{18}\text{O}$ and $^{238}\text{U} + ^{22}\text{Ne}$ reactions	166
5.6	Gaussian fit to the experimental ²⁴⁸ Cm + ¹⁵ N reaction	167

5.7	1n-exit channel systematics	171
5.8	2n-exit channel systematics	172
5.9	4n-exit channel systematics	174
5.10	New HIVAP predictions for the ²⁰⁸ Pb(⁴⁸ Ca,xn) ^{256-x} No reaction	177
5.11	New HIVAP predictions for the ²⁰⁸ Pb(⁵⁰ Ti,xn) ^{258-x} Rf reaction	178
5.12	New HIVAP predictions for the ²⁰⁹ Bi(⁵⁰ Ti,xn) ^{259-x} Db reaction	179
5.13	New HIVAP predictions for the ²⁰⁹ Bi(⁵¹ V,xn) ^{260-x} Sg reaction	180
5.14	Deformation energy versus length	185
5.15	Probability of compound nucleus formation versus injection point length	186

List of Tables

1.1	IUPAC recommended names and symbols for elements 101-109	5
1.2	Calculated E^*_{min} values for the discovery experiments	18
2.1	Reisdorf and Schädel parameters for the HIVAP code	32
3.1	BGS reaction specifics	50
3.2	Event word list for the ²⁰⁹ Bi(⁵¹ V,2n) ²⁵⁸ Sg experiment	56
4.1	Summary of literature half-lives, branching ratios, and primary α -energies	60
4.2	Alpha peak assignments for the ²⁴⁸ Cm(¹⁵ N,3n) ²⁶⁰ Lr experiment	65
4.3	Decay tables for the ²⁴⁸ Cm(¹⁵ N,3n) ²⁶⁰ Lr experiment	67
4.4	MLDS results for the ²⁴⁸ Cm(¹⁵ N,3n) ²⁶⁰ Lr experiment	69
4.5	Reaction specifics for the ²³⁸ U(¹⁸ O,4n) ²⁵² Fm experiment	77
4.6	Alpha peak assignments for the ²³⁸ U(¹⁸ O,4n) ²⁵² Fm experiment	81
4.7	Decay tables for the ²³⁸ U(¹⁸ O,4n) ²⁵² Fm experiment	83
4.8	MLDS results for the ²³⁸ U(¹⁸ O,4n) ²⁵² Fm experiment	84
4.9	Reaction specifics for the ²³⁸ U(²² Ne,xn) ^{260-x} No experiment	88
4.10	MLDS results for the ²³⁸ U(²² Ne,4n) ²⁵⁶ No experiment	100
4.11	Reaction specifics for the ²⁰⁸ Pb(⁴⁸ Ca,xn) ^{256-x} No experiment	103
4.12	Decay tables for the ²⁵⁵ No and ²⁵⁴ No experiments	110
4.13	MLDS results for ²⁵⁵ No and ²⁵⁴ No	111
4.14	Production cross sections for ²⁵⁵ No and ²⁵⁴ No	113
4.15	Decay tables and MLDS results for the ²⁵³ No and ²⁵² No experiments	118
4.16	Reaction specifics for the ²⁰⁸ Pb(⁵⁰ Ti,xn) ^{258-x} Rf experiment	123
4.17	Reaction specifics for the ²⁰⁹ Bi(⁵¹ V,2n) ²⁵⁸ Sg experiment	147

4.18	Correlation summary of the ²⁰⁹ Bi(⁵¹ V,2n) ²⁵⁸ Sg reaction	149
5.1	Summary of experimental hot fusion production cross sections	154
5.2	Summary of experimental cold fusion production cross sections	155
5.3	Results of gaussian fits to the GSI data	158
5.4	Results of gaussian fits to the experimental cold fusion data	163
5.5	Results of gaussian fits to the experimental hot fusion data	168

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1 Introduction

1.1 Discovery of the transuranium elements

Chemistry has played an important role in the discovery and positive identification of new artificial elements since the production and identification of neptunium and plutonium in 1940-41. In 1872, Mendeleev furthered the progress for the search of new elements with his formulation of the periodic table. This helped give insight on where to look for new elements, and how these new elements might behave chemically. In the 1930's, technical advances and the invention of the cyclotron by E. O. Lawrence, opened up a new era in element discovery.

The first elements heavier than uranium, neptunium (Z = 93) and plutonium (Z = 94), were produced and identified in 1940 and 1941. The discoveries involved the irradiation of uranium by neutrons and deuterium ions and identification through chemistry. Neptunium was discovered by E. M. McMillian in the neutron activation of uranium [McM1940] in which a single neutron was added to a 238 U nucleus, producing 239 U which then beta-decayed to neptunium.

$${}^{238}_{92}U({}^{1}_{0}n,\gamma){}^{239}_{92}U \xrightarrow{\beta^{-}}{}^{23.5}_{93}Np$$
 (1.1)

Plutonium was discovered not long afterward through the interaction of deuterium with uranium [Sea1946]. Neptunium was the reaction product which then beta-decayed to plutonium.

$${}^{238}_{92}U({}^{2}_{1}H,2{}^{1}_{0}n){}^{238}_{93}Np \xrightarrow{\beta^{-}}_{2.12d} \xrightarrow{238}^{238}Pu$$
 (1.2)

Americium (Z = 95) and curium (Z = 96) were the next two heavy elements to be discovered. Initial experiments to chemically separate and identify them as homologues

of iridium and platinum were unsuccessful until Glenn T. Seaborg proposed a new actinide series similar to the lanthanide series [Sea1945]. Curium was identified first through the irradiation of a long-lived isotope of plutonium with alpha-particles [Sea1945].

$${}^{239}_{94}Pu\left({}^{4}_{2}He, {}^{1}_{0}n\right){}^{242}_{96}Cm \tag{1.3}$$

Chemical separation as the trivalent species later confirmed the production of the new element. Americium was later identified in the multiple neutron capture of the same long-lived plutonium isotope [Ghi1950].

$${}^{239}_{94}Pu\left(2_{0}^{1}n,\gamma\right){}^{241}_{94}Pu \xrightarrow{\beta^{-}}_{14.4y} \xrightarrow{}^{241}_{95}Am \tag{1.4}$$

Seaborg formally proposed the actinide concept in the discovery letter for curium, supported by the separation of curium and americium in the trivalent oxidation state and with it a new periodic table [Sea1945]. This postulation was extremely important to the discovery of new heavier elements. Seaborg's proposal paved the way for chemical identification of the heavier elements by predicting that the actinides behaved chemically similarly to their homologue lanthanides. The lanthanide series filled the inner 4f shell, while the actinides fill the 5f shell [Hff1999]. Based on this concept, new heavy elements would fill the actinide row up to element 103. These new actinides were predicted to all exhibit stable 3+ oxidation states, except for element 102, which was predicted to have a stable 2+ oxidation state like its homologue ytterbium. Chemical separations used at the time involved changing the oxidation state of the species of interest while not changing the oxidation states of the impurities, and then extracting the species of interest effectively separating it from the impurities. With the majority of

these new actinides exhibiting a stable 3+ oxidation state, new chemical separation techniques were needed.

With the actinide concept in mind, work began on the discovery of still heavier elements. The technology of the time was limited to light ion beams, so heavier target material needed to be produced in significant quantities to produce a target suitable for irradiations. In 1949, berkelium was discovered through the irradiation of americium with helium ions [Tho1950a].

$${}^{241}_{95}Am\left({}^{4}_{2}He,2{}^{1}_{0}n\right){}^{243}_{97}Bk\tag{1.5}$$

Positive identification of these new activities as isotopes of new elements required new chemical separation techniques. Elution from ion-exchange resin columns with different elutants was performed to separate species with similar oxidation states. These new techniques were used in the positive identification of berkelium and californium.

Californium was produced in 1950 through the irradiation of curium with helium ions [Tho1950b].

$${}^{242}_{96}Cm\left({}^{4}_{2}He, {}^{1}_{0}n\right){}^{245}_{98}Cf \tag{1.6}$$

Chemical separations of californium from the other actinides produced in this reaction were critical to its discovery. A cation exchange column technique was used to elute californium before the similar trivalent actinides berkelium and curium [Tho1950b].

The discovery of elements 99 and 100 did not occur through the use of accelerated projectiles and a target. Great effort was given at the time to use the heaviest and most neutron-rich targets available. However, short target material half-lives, difficulty in the production of significant quantities of target material, and the limited

number of beam options slowed the continuing search for these two new elements. Einsteinium and fermium, elements 99 and 100 were discovered in the debris from the "Mike" thermonuclear device tested in the South Pacific on November 1, 1952 [Ghi1955a]. Early chemical separations from the "Mike" test debris identified the heaviest known isotopes of plutonium, ²⁴⁶Pu and ²⁴⁴Pu. This provided evidence that the ²³⁸U used in the "Mike" test had captured at least eight neutrons forming ²⁴⁶U which then decayed to ²⁴⁶Pu via successive beta decay. This led the researchers to speculate that more neutrons might have been captured and maybe heavier elements could be detected from the multiple beta-decays that followed the multiple neutron captures. This indeed proved to be true as an intensive search for elements 99 and 100 by researchers at the Radiation Laboratory in Berkeley, Argonne National Laboratory in Illinois, and the Los Alamos Scientific Laboratory verified the multiple neutron capture of ²³⁸U. These researchers performed the first chemical separation and isolation of elements 99 and 100 from debris recovered from the "Mike" test. Chemical separation techniques similar to those used to discover curium, berkelium and californium were used. Elution from cation exchange resins showed einsteinium and fermium in their predicted positions.

Names for the previous elements were all proposed by their discoverers soon after they were identified as being new elements. In 1997, the International Union of Pure and Applied Chemistry (IUPAC) assigned credit for discovery and approved the names of elements 101-109. The elements name and symbol can be found in Table 1.1 [Jup1997].

Table 1.1: IUPAC approved names and symbols for elements 101-109 [Iup1997].

Element	Name	Symbol	
101	mendelevium	Md	
102	nobelium	No	
103	lawrencium	Lr	
104	rutherfordium	Rf	
105	dubnium (hahnium*)	Db (Ha*)	
106	seaborgium	Sg	
107	bohrium	Bh	
108	hassium	Hs	
109	meitnerium	Mt	

^{*}Hahnium and the symbol "Ha" appeared in the literature for chemical studies of element 105 prior to 1997.

In 1955, mendelevium was discovered in the bombardment of einsteinium atoms by helium ions. Enough ²⁵³Es had been separated to form a target, which was then irradiated by helium ions [Ghi1955b].

$${}^{253}_{99}Es\left({}^{4}_{2}He, {}^{1}_{0}n\right){}^{256}101 \tag{1.7}$$

This experiment was important because mendelevium was the first heavy element produced and identified using atom-at-a-time techniques which became important in future experiments. It would also incorporate a new technique to simplify the chemical analysis. The new technique used a catcher foil placed directly behind the target to collect the recoiling products as they left the target. This too would become important in future research. It was also the last experiment performed using light ion beams ($Z \le 2$, $A \le 4$). Beams heavier than helium were required to produce the elements with Z > 101.

Nobelium, element 102 was erroneously reported discovered in 1957. Irradiations were performed in the cyclotron at the Nobel Institute in Stockholm, Sweden. Curium targets were irradiated by carbon ions at various energies. It was believed that the following reactions were observed [Fie1957],

$${}^{244}_{96}Cm\left({}^{13}_{6}C,6{}^{1}_{0}n\right){}^{251}102\tag{1.8}$$

$${}^{244}_{96}Cm\left({}^{13}_{6}C,4{}^{1}_{0}n\right){}^{253}102\tag{1.9}$$

Chemical separations were performed to isolate nobelium from the other reaction products. Separations were performed under the assumption that nobelium behaved similarly to mendelevium, fermium and einsteinium, exhibiting a stable 3+ oxidation state in aqueous solution. This was later proven incorrect when nobelium was shown to

exhibit a stable 2+ oxidation state in aqueous solution [Mal1968]. Soon afterwards, additional experiments to find nobelium were performed in Berkeley. Ghiorso and coworkers performed similar experiments with carbon on curium [Ghi1958].

$${}^{246}_{96}Cm\left({}^{12}_{6}C,4{}^{1}_{0}n\right){}^{254}102\tag{1.10}$$

Researchers at Berkeley identified nobelium through the chemical identification of its alpha-decay daughter ²⁵⁰Fm and separately through the direct counting of ²⁵²No collected in a catcher foil [Hff1998]. Work was also performed at the same time at Dubna using plutonium targets [Fle1958].

$${}^{239}_{94}Pu\left({}^{16}_{8}O,xn\right){}^{255-x}102\tag{1.11}$$

Through the efforts of researchers in Berkeley and in Dubna, the discovery of nobelium was possible. A more through discussion of the events of the discoveries of nobelium and the rest of the heavy elements can be found in [Sea1990] and [Hff1998, Hff2000]. This was the first of the discovery experiments that used a beam heavier than helium. Four neutrons were evaporated in this reaction because of the large excitation energy of the resulting compound nucleus.

Lawrencium, element 103, and the last of the actinides, was discovered in 1961. Californium targets were bombarded by intense beams of accelerated boron ions to produce various isotopes of lawrencium [Ghi1961].

$${}^{249-252}_{98}Cf\left({}^{10,11}_{5}B,4_{0}^{1}n\right){}^{255-259}103\tag{1.12}$$

Lawrencium was detected by collecting the activity from the reaction on a tape which was moved between a series of alpha-particle detectors [Sea1990]. Various isotopes

were observed by examining the resulting spectra of alpha-particle energies. From the information gathered, the most probable isotope produced was found to be ²⁵⁷Lr. It is the product resulting from the emission of three, four and five neutrons from the fusion of ¹⁰B with ²⁵⁰⁻²⁵²Cf, respectively, and four, five and six neutrons from the fusion of ¹¹B with ²⁵⁰⁻²⁵²Cf, respectively [Ghi1961]. This behavior is consistent with the reactions that produce large amounts (~ 40 MeV) of excitation energy in the compound nucleus. Such reactions that produce compound nuclei from reactions of ions with actinide targets are called "hot fusion" reactions. Increased excitation energy leads to a smaller probability of surviving from fission due to increased neutron evaporation steps required to remove the excess energy. With the fission probability significantly higher than neutron emission probability at each step, the total fission survivability probability is small. Hot fusion reactions have higher compound nucleus fusion probabilities though as incident projectile energies are well above the interaction barrier.

Enhanced physical detection techniques [Ghi1967a, Ghi1967b] and enhanced beam accelerators (HILAC and SuperHILAC) were used to discover the transactinide elements 104, 105 and 106. The californium target so successfully used to produce lawrencium was used again with various beams to produce the new heavy elements 104, 105, and 106, later named rutherfordium, dubnium and seaborgium [Jup1997]. Rutherfordium was produced in the hot fusion reaction of carbon ions with californium [Ghi1969].

$${}^{249}_{98}Cf\left({}^{12}_{6}C,4{}^{1}_{0}n\right){}^{257}104\tag{1.13}$$

Dubnium was produced a short time later in the reaction of californium with nitrogen ions [Ghi1970].

$${}^{249}_{98}Cf\left({}^{15}_{7}N,4{}^{1}_{0}n\right){}^{260}105\tag{1.14}$$

Seaborgium was produced in the reaction of californium with oxygen ions [Ghi1974].

$${}^{249}_{98}Cf\left({}^{18}_{8}O,4{}^{1}_{0}n\right){}^{263}106\tag{1.15}$$

The highlights of the discoveries of elements 104-106 as well as additional discovery claims can be found in Seaborg and Hoffman [Sea1990, Hff1998, Hff2000]. All of these experiments used physical means to verify the discovery of the new elements. The most reliable physical method was to observe the alpha decay of the isotopes produced and link them to the alpha decay of previously known isotopes. This method was used for the discovery of rutherfordium, dubnium and seaborgium. For example, ²⁶³Sg was linked genetically to ²⁵⁹Rf which was genetically linked to ²⁵⁵No. Detection systems were designed to provide optimum conditions for detecting these genetic relationships. The chemical properties of these transactinide elements were predicted to be similar to their d-block homologues. Chemical separations to test these ideas were not performed until production rates were increased and chemical separation techniques became more sophisticated. Various production methods were also examined in the hope of producing isotopes of these heavy elements with long enough half-lives to perform chemistry. The first chemistry experiments on rutherfordium [Zva1969], dubnium [Zva1970] and seaborgium [Shä1997] were performed in 1969, 1970 and 1997, respectively.

After the discovery of seaborgium, it would again be sometime before the discovery of a new heavy element. Extremely low production rates, small cross sections,

and limited beam currents all contributed to the difficulty in producing new elements. All of the approaches to this point involved the use of actinide targets to produce the heavy elements. In 1975, Oganessian and co-workers [Oga1975] postulated that through the use of lead and bismuth targets and beams with $Z \sim 20$, compound nuclei could be created with smaller excitation energies which would increase the survivability of the evaporation residues from fission. This new reaction was called a "cold fusion" reaction, as the compound nucleus was "colder" than those produced in hot fusion reactions. The cold fusion reactions targets ²⁰⁸Pb and ²⁰⁹Bi have binding energies (²⁰⁸Pb: 1636 MeV, 209 Bi: 1640 MeV), that are 150 MeV – 200 MeV smaller than the binding energies of hot fusion targets like ²³⁸U (1802 MeV) and ²⁴⁹Cf (1863 MeV). In addition the compound nucleus Q-value is approximately 100 MeV smaller for cold fusion reactions than hot fusion reactions. For example, the O-value for ²⁵⁸Rf, the compound nucleus from the 208 Pb + 50 Ti reaction is –169 MeV, whereas the Q-value for 262 Rf, the compound nucleus from the ²³⁸U + ²⁴Mg reaction is –69 MeV. These two factors lead to smaller compound nucleus excitation energies in cold fusion reactions.

In 1981, element 107 was produced in the cold fusion reaction of bismuth with chromium ions [Mün1981].

$${}^{209}_{83}Bi\left({}^{54}_{24}Cr, {}^{1}_{0}n\right){}^{262}107 \tag{1.16}$$

This was the first cold fusion reaction used in the discovery of a new heavy element. The evaporation of only one neutron is characteristic of smaller excitation energies in the compound nucleus. This discovery would not have been possible without the construction of the UNILAC accelerator and SHIP (Separator for Heavy Ion reaction

Products) velocity filter [Mün1979] under the direction of Armbruster at GSI in Darmstadt.

Following the successful discovery of bohrium, elements 108 and 109 were quickly discovered in 1982-84 [Mün1984a, Mün1987, Mün1982, Mün1984b] using cold fusion reactions of heavy ions with ²⁰⁸Pb or ²⁰⁹Bi targets.

$${}^{208}_{82}Pb\left({}^{58}_{26}Fe, {}^{1}_{0}n\right){}^{265}108\tag{1.17}$$

$${}^{209}_{83}Bi\left({}^{58}_{26}Fe, {}^{1}_{0}n\right){}^{266}109\tag{1.18}$$

Elements 110 – 112 were not discovered by the GSI team until nearly ten years later after improvements were made in the efficiency and detection system of SHIP and the use of more intense beams [Hof1995a, Hof1995b, Hof1996].

$${}^{208}_{82}Pb\left({}^{62}_{28}Ni,{}^{1}_{0}n\right){}^{269}110\tag{1.19}$$

$${}^{209}_{83}Bi\left({}^{64}_{28}Ni, {}^{1}_{0}n\right){}^{272}111\tag{1.20}$$

$${}^{208}_{82}Pb\left({}^{70}_{30}Zn,{}^{1}_{0}n\right){}^{277}112\tag{1.21}$$

The reactions producing elements 111 and 112 have not yet been confirmed outside of GSI [Hof2001]. Additional information on the discoveries of elements 107 through 110 can be found in [Sea1990] and of elements 107 through 112 in [Hff1998, Hff2000].

The problem with using cold fusion reactions to extend the discovery to heavier elements (Z > 112) resides in the fact that production cross sections for the best reactions are approximately 1 picobarn (10^{-36} cm²). The current sensitivity of detection equipment is approximately 1 picobarn as well. It can be seen that the best reactions to produce

these heavy elements have cross sections that decrease rapidly with Z, and increasingly better sensitivities are required.

To create the superheavy elements (Z > 112), detection equipment needs to be improved, beam currents need to be increased, or a completely different route needs to be investigated. The possibility of using actinide targets with the same heavy ion beams used in cold fusion reactions to produce elements heavier than 112 was discussed as early as 1981 [Oga1981]. Reactions of actinide targets and intense ⁴⁸Ca beams produce compound nucleus Q-values similar to the compound nucleus Q-values of the cold fusion reactions, leading to slightly larger compound nucleus excitation energies.

As a result of better equipment and increased detection sensitivity as well as the ability to produce rather intense ⁴⁸Ca beams, elements 114 and 116 were reported produced. In 1999, element 114 was reported in the reaction of ⁴⁸Ca ions with neutron-rich plutonium targets [Oga1999a, Oga2000a].

$${}^{244}_{94}Pu\left({}^{48}_{20}Ca, 4{}^{1}_{0}n\right){}^{288}114\tag{1.22}$$

Built upon the success of the reactions of ⁴⁸Ca on plutonium and uranium reported for the production of neutron-rich element 112 isotopes [Oga1999b], high intensity ⁴⁸Ca beams were used to irradiate ²⁴⁸Cm targets to produce element 116 [Oga2000b].

$${}^{248}_{96}Cm\left({}^{48}_{20}Ca, 4{}^{1}_{0}n\right){}^{292}116\tag{1.23}$$

Only five atoms of element 114 [Oga2001a, Oga2000a, Oga1999a] and one atom of element 116 [Oga2000b] have been produced and these experiments have not been confirmed. The expectation of the emission of four neutrons is consistent with compound

nucleus excitation energies between 30 and 40 MeV and the classification of these reactions as hot fusion reactions.

The search for still heavier elements will continue to use both cold and hot fusion reactions. Advanced beam development will lead to higher intensity beams which will require innovative target designs to dissipate the heat produced. Separation devices will be improved to detect activity faster and more accurately than before. Aside from the instrumentation aspect of the search, it is also important to understand the physical reasons for the success of these individual reaction paths, cold- and hot fusion, and when to use one type of reaction over another.

1.2 Cold and hot fusion

The initial predictions on cold fusion reactions [Oga1975] was based on a simple compound nucleus excitation energy calculation. The calculation showed a minimum excitation energy (E^*_{min}) in the compound nucleus formed in reactions of projectiles with masses around 45. The calculation is based on finding the excitation energy of the compound nucleus at the interaction barrier (B_{int}). The interaction barrier in this calculation is a Coulomb potential barrier (V_{coul}).

$$E_{\min}^* = B_{\inf} + Q, \tag{1.24}$$

$$B_{\rm int} = V_{coul} \tag{1.25}$$

$$V_{coul} = \frac{Z_1 Z_2 e^2}{r_e \left(A_1^{\frac{1}{3}} + A_2^{\frac{1}{3}} \right)}$$
(1.26)

$$Q = (M_1 + M_2 - M_{CN})c^2 (1.27)$$

The effective interaction radius (r_e) was taken as 1.45 fm and the nuclear masses were taken from Myers and Swiatecki [Mye1966]. Using this simple calculation and various projectile and target combinations for the production of fermium and rutherfordium, Oganessian and co-workers produced the graph seen in Figure 1.1. It is easy to notice from Figure 1.1 that the minimum of this curve appears around a projectile mass of 45. This meant that ion beams like ⁴⁰Ar and ⁴⁸Ca with ²⁰⁸Pb and ²⁰⁹Bi targets might be more effective in forming the heavy elements than actinide target reactions with lighter ions. This can be seen in Figure 1.2.

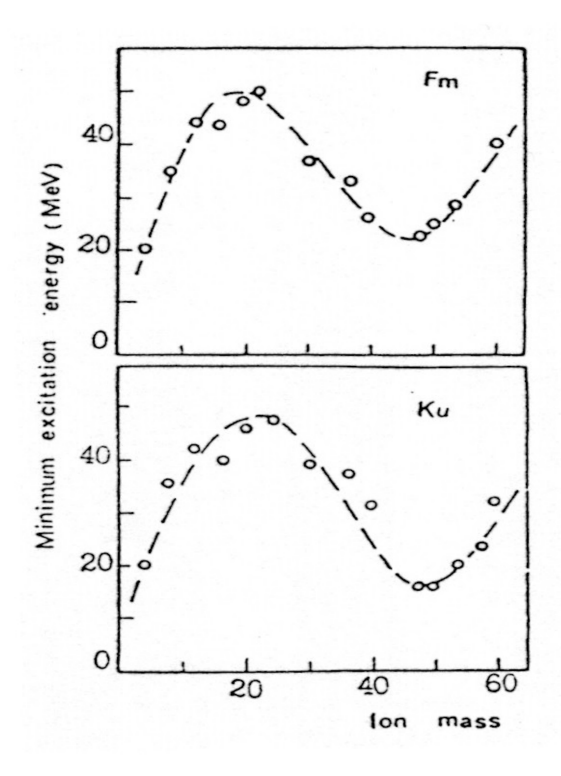


Figure 1.1: Graphs of projectile (ion) mass versus E^*_{min} for various projectile-target reactions leading to 248 Fm and 258 Rf (denoted Ku). The dashed curves are drawn through the calculated E^*_{min} values shown by the points [Oga1975].

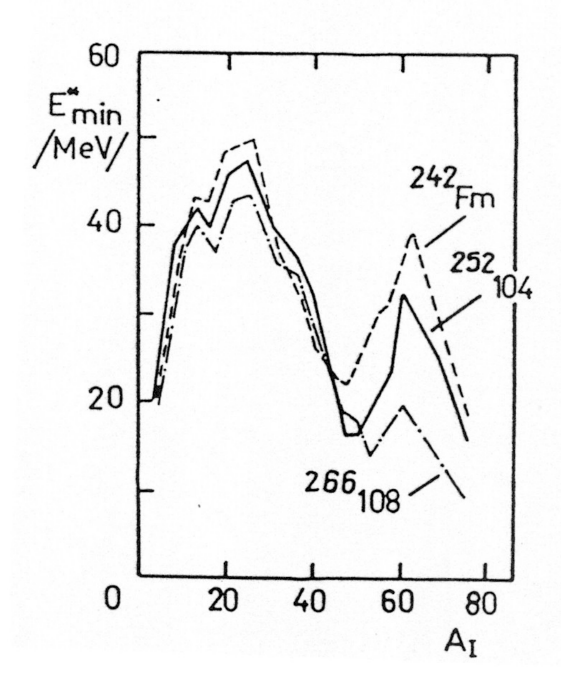


Figure 1.2: The minimum excitation energy versus the projectile mass A_I for different target-projectile combinations leading to 242 Fm, 252 Rf and 266 Hs compound nuclei [Oga1981].

One can see from Figure 1.2 that the trend continues for even heavier elements like hassium (element 108). The minimum in these curves does shift to higher masses with the production of the heavier elements, meaning that the corresponding target is centered around lead. This postulate led to the discovery of elements $107 \le Z \le 112$ by researchers at GSI in Darmstadt using their velocity filter SHIP [Mün1979]. Table 1.2 illustrates the E^*_{min} values for the confirmed discovery reactions from mendelevium through element 110 and reported discovery reactions for elements 111, 112, 114 and 116.

Table 1.2 shows that the minimum excitation for the discovery of elements 101 through 106 remained constant around 40 MeV. Hot fusion experiments to produce elements heavier than seaborgium were difficult due to lower fusion probabilities. Lower fusion probabilities combined with constant high fission competition in the exit channel reduced the production cross sections for hot fusion reactions. The use of cold fusion reactions helped solve the decreasing production cross section problem. Smaller compound nucleus excitation energies meant reduced fission competition in the exit channel and higher production cross sections. Only small gains were made as decreases in the fusion probability for cold fusion reactions continued the decline of production cross sections. Figure 1.3 illustrates the decrease in cross section of various 1n-exit channels for cold fusion reactions for elements 102 – 113 [Hof2000].

Table 1.2: Projectile, target and compound nucleus Z and A for the discovery reactions for elements 101 –110 and reported reactions for elements 111,112,114, and 116 and calculated E^*_{min} values. Z_p , Z_t , and Z_{CN} refer to the atomic number of the projectile, target, and compound nucleus. A_p , A_t , and A_{CN} refer to the atomic mass number of the projectile, target, and compound nucleus.

Z_p	A_p	Z _t	A _t	Z _{CN}	A _{CN}	E* _{min} (MeV) ^a
2	4	99	253	101	257	17.48
6	12	96	244	102	256	37.72
5	10	98	251	103	261	44.46
6	13	98	249	104	262	38.45
7	15	98	249	105	264	38.63
8	18	98	249	106	267	40.19
24	54	83	209	107	263	14.31
26	58	82	208	108	266	11.49
26	58	83	209	109	267	10.83
28	62	82	208	110	270	8.05
28	64	83	209	111	273	4.13
30	70	82	208	112	277	-0.78
20	48	94	244	114	292	19.11
20	48	96	248	116	296	18.14

 $^{^{\}text{a}}$ An r_{e} value of 1.45 fm and masses from [Lir1976, Lir2001] were used.

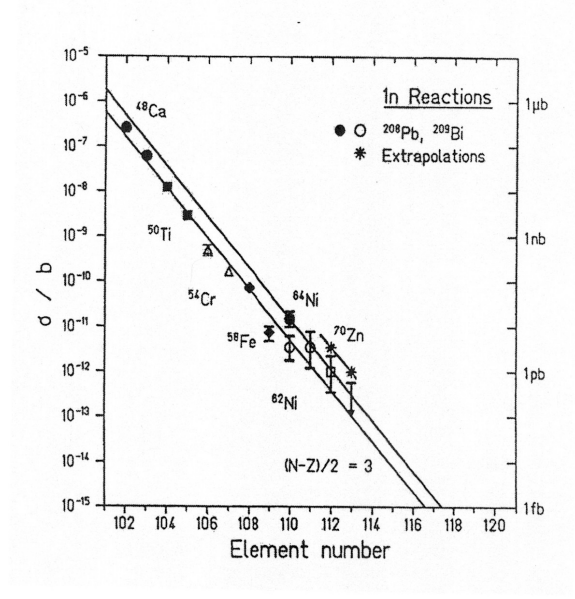


Figure 1.3: 1n-exit channel cross sections for elements 102-113 from cold fusion reactions of various projectiles with lead and bismuth targets [Hof2000].

The deciding factor in choosing between hot and cold fusion reactions is the desired end product. Currently hot fusion reactions using ⁴⁸Ca projectiles and actinide targets appear to be the route to the superheavy elements because of the neutron-rich character of the heavy element isotopes produced. Cold fusion reactions with ²⁰⁸Pb and ²⁰⁹Bi targets are preferred for the study of the neutron-deficient transactinide isotopes.

One advantage to using hot fusion reactions is the ability to form compound nuclei that are neutron-rich and have longer half-lives. Trial calculations have predicted a doubly deformed shell around Z=108 and N=162 [Pat1989, Pat1991]. Hot fusion reactions are a possible reaction mechanism for the formation of such neutron-rich nuclides. A second advantage to using the hot fusion mechanism is the enhanced probability of fusion of the projectile-target system [Oga1994]. The highest production cross sections in hot fusion reactions occur in the 4n- and 5n- exit channels which are located well above the projectile-target interaction barrier. The disadvantages to hot fusion reactions are all due to the large amount of excess excitation energy in the compound nucleus. The higher excitation energy destabilizes the compound nucleus by eliminating the shell effects present in the de-excited evaporation residues [Fle1976]. Numerous neutron evaporation steps are required to evaporate this excess excitation energy leading to an enhanced fission probability, and therefore lower production cross sections.

The main advantage of using cold fusion reactions is the enhanced survivability to fission during compound nucleus de-excitation. With smaller compound nucleus excitation energies, less neutron evaporation steps are required leading to a smaller fission probability. This smaller probability to fission is also related to the fact that the

smaller excitation energy does not tend to eliminate the shell effects. The remaining shell effects add stability to the compound nucleus reducing the chance of fission [Oga1994]. Additionally, fusion of more symmetrical projectile-target combinations leads to cooler compound nuclei [Arm1985]. Another advantage to the cold fusion reaction mechanism is the ability of these reactions to produce neutron-deficient nuclei. This gives access to the study of the decay properties and lifetimes of nuclei away from areas of spherical and deformed stability. Cold fusion reactions, however, do not produce neutron-rich nuclei, and therefore cannot form the neutron-rich superheavy elements. For example, a cold fusion reaction to form rutherfordium, ²⁰⁸Pb + ⁵⁰Ti, forms the compound nucleus, ²⁵⁸Rf, with only 154 neutrons whereas a hot fusion reaction to form rutherfordium, ²³⁸U + ²⁶Mg, forms the compound nucleus, ²⁶⁴Rf, with 160 neutrons.

The following three figures illustrate the decrease in cross sections for hot and cold fusion reactions as a function of increasing atomic number. Figure 1.4 compares ²⁰⁸Pb and ²⁰⁹Bi cold fusion reactions with ²³⁸U and ²⁴⁴Pu target based hot fusion reactions [Oga2001b]. Figure 1.5 displays additional hot and cold fusion cross sections as well as some reported experimental work on the production cross sections of elements with Z > 112 [Mün2001]. Finally, Figure 1.6 illustrates for a variety of reactions, the maximum cross section recorded for a given element from fermium through 116 produced in either hot and cold fusion reactions [Ghi1961, Oga1975, Mün1982, Mün1984, Hof1995a, Hof1995b, Hof1996, Oga1999a, Oga2000a, Shi1986, Sik1968, Gäg1989, Nit1981, Heβ1997, Kra1992, Heβ2001a, Gre1994, Mün1985, Wil2000, Mün1989, Tür2001, Hof1998, Hof2001, Oga1999b, Laz1996, Moo2001].

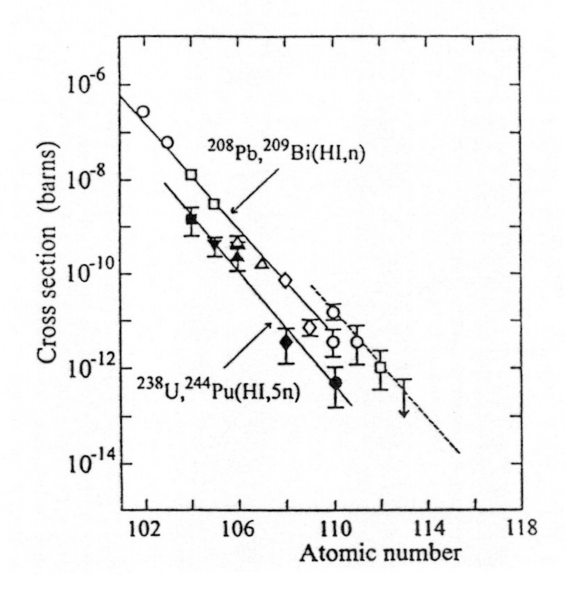


Figure 1.4: Production cross section versus atomic number for various cold and hot fusion reactions [Oga2001b]. HI denotes heavy ion projectile.

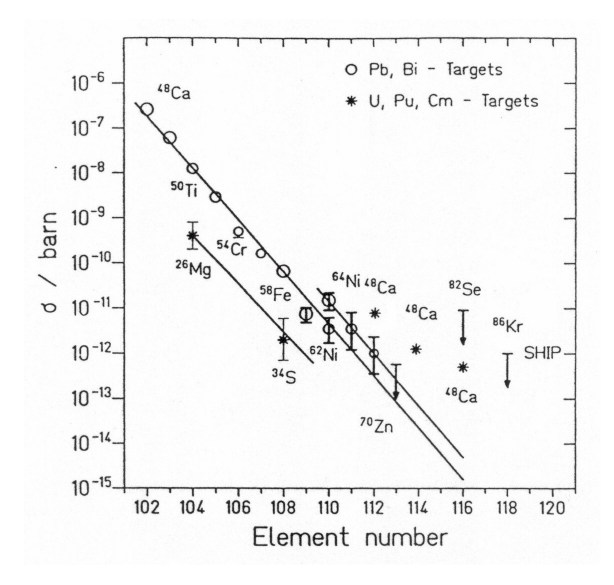


Figure 1.5: Cross section versus increasing element number for cold fusion reactions with lead and bismuth targets and hot fusion reactions with uranium, plutonium and curium targets. Recent work on elements $Z \ge 116$ is also included. Adapted from [Mün2001].

Cold Fusion/Hot Fusion Comparison

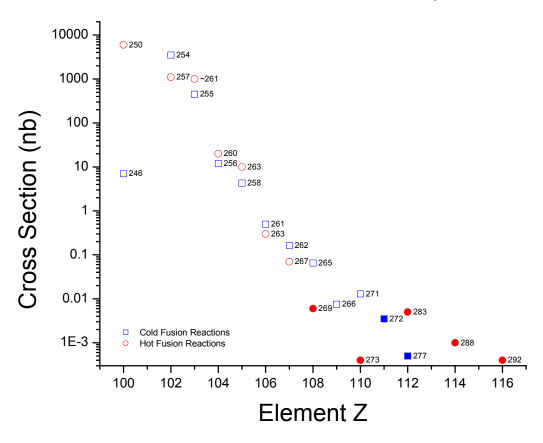


Figure 1.6: Cross section versus Z. The maximum cross section for a given reaction (cold/hot fusion) is shown. Hot fusion in circles and cold fusion in squares. Open symbols represent confirmed reactions while solid symbols represent unconfirmed reactions. The mass number for the isotope produced is given to the left or right of each symbol [Ghi1961, Oga1975, Mün1982, Mün1984, Hof1995a, Hof1995b, Hof1996, Oga1999a, Oga2000a, Shi1986, Sik1968, Gäg1989, Nit1981, Heβ1997, Kra1992, Heβ2001a, Gre1994, Mün1985, Wil2000, Mün1989, Tür2001, Hof1998, Hof2001, Oga1999b, Laz1996, Moo2001].

From Figure 1.6 it can be seen that the hot fusion reactions do indeed produce nuclei that are more neutron-rich. Hot fusion reactions also produce evaporation residues with higher cross sections for the elements $Z \le 105$ and $Z \ge 112$. The lone exception is the production of nobelium using the cold fusion of doubly magic ⁴⁸Ca and ²⁰⁸Pb.

The study of hot and cold fusion reactions can be extremely enlightening in terms of the production routes to the heavy elements. The importance of understanding hot and cold fusion does not end with the heavy elements. Understanding which method is preferred over the other for a given region of nuclides can facilitate the study of a particular area of the chart of nuclides in greater detail. By examining particular reactions and making subtle changes in the choice of target and projectile combinations, a greater understanding of the structure of the nuclei in the region of the heaviest elements can be developed.

1.3 Scope

In this dissertation, experimental investigations of both hot and cold fusion reactions are described. Three hot fusion reactions will be examined:

248Cm(15N,xn)263-xLr, 238U(18O,4n)252Fm, and 238U(22Ne,xn)260-xNo. The 3n- and/or 4n-exit channel was studied in each experiment. Five cold fusion experiments are examined:

208Pb(48Ca,xn)256-xNo, 208Pb(50Ti,xn)258-xRf, 208Pb(51V,2n)257Db, 209Bi(50Ti,xn)259-xDb, and

209Bi(51V,2n)258Sg. A review of three of the various computer codes available for the prediction of production cross sections are reviewed. An examination of the results from various hot and cold fusion experiments is presented and compared with literature values. The relevance of these experiments to the development of a better understanding of hot and cold fusion excitation functions is discussed. The importance of projectile odd-even effects and target odd-even effects in the production cross sections of neutron deficient nuclides is discussed. The relevance of computer codes in calculating cross sections, and how these codes can help in understanding the physics behind the production of nuclides of the heaviest elements is pointed out.

2 Evaporation codes

2.1 JORPLE and SPIT

Cross sections are important to the study of hot and cold fusion reactions. A cross section is defined as a measure of the probability or likelihood that a specific reaction will occur. Different reactions leading to the same isotope will have different cross sections. These cross sections can be compared to evaluate the best mechanism for the synthesis of that particular isotope. Cross sections are measured in units of area called barns (b) where one barn is equal to 10^{-24} cm².

Heavy element experimental cross section measurements are performed on isotopes that have small (< 1 millibarn) cross sections. This means that the amount of time required for the experiment can be quite long, and therefore expensive in terms of equipment costs and beam time. Knowing the expected cross section can greatly reduce the amount of time spent on an experiment by helping determine what experimental energies should be tested and for how long.

The foundations of the JORPLE code are based in the estimations of the production cross sections for heavy element reactions [Sik1966]. The computer code was written in 1970 [Alo1970, Alo1974]. This code provides a general prediction of the cross sections that could be expected from a variety of heavy element reactions.

The interaction potential for the JORPLE code is based on a sum of individual potentials [IIj1982], a Coulomb term like Equation 1.26, a rotational term, and a nuclear term. The nuclear part of the equation resembles the Woods-Saxon potential but is partially modified [Alo1974] by adding in terms dependant on the orientation angles of the nuclear deformation axis with respect to the incident beam direction. The total fusion

probability in the JORPLE code is solely based on the barrier penetration probability, which is calculated using the Hill and Wheeler approximation [Hill953].

The de-excitation of the compound nucleus is performed using the Jackson model to calculate neutron evaporation probabilities [Sik1968a, Sik1968b]. The values for the (Γ_n/Γ_f) values are calculated according to Sikkeland and co-workers [Sik1968a],

$$\log\left(\frac{\Gamma_n}{\Gamma_f}\right) = -0.276Z + \begin{cases} 5.46 + 0.140N, N \le 153\\ 19.23 + 0.050N, N \ge 153 \end{cases}$$
(2.1)

where Z and N refer to the atomic number and neutron number of the compound nucleus.

By combining the effects of all of the partial angular momentum waves through the interaction barrier, and adding in the barrier penetration factor with a compound nucleus de-excitation factor, the JORPLE code is a simple calculation that gives relatively good production cross section estimations.

The SPIT code is a modification of the JORPLE code developed to increase the accuracy of cross section predictions [Wil1988]. Modifications were made to the original code by changing the interaction potential in the attempt to make the estimations better reflect the existing experimental data.

The SPIT code differs from the JORPLE code in using a different Coulomb potential as well as a different nuclear potential. The rotation part of the interaction potential is kept the same as in the JORPLE code. The Coulomb potential is taken from Bondorf, Sobel and Sperber [Bon1974]. The nuclear part of the interaction potential is based on the Bass proximity potential [Bas1977].

Taking the new nuclear and Coulomb parts to the interaction potential, the first step in the SPIT code is the determination of the barrier energy. From here, using the same calculations as in the JORPLE code, the barrier penetration factor and the compound nucleus cross section are calculated. From there, the neutron evaporation probability and neutron width to fission width ratio are calculated and multiplied together with the compound nucleus cross section to give the production cross section for a given energy and number of neutrons evaporated.

Using these modified Coulomb and nuclear potentials leads to enhanced accuracy in predicting production cross sections for the heavy elements. The SPIT and JORPLE codes and their ability to predict production cross sections for heavy ion projectile actinide target reactions have been reviewed in [Hay1988, Moo1990]. In most cases for reactions involving projectiles equal to or lighter than oxygen, the SPIT code can reproduce the experimental cross section within an order of magnitude or two. Heavier projectile beams lead to larger discrepancies. As new reaction mechanisms were developed to study heavier element systems, a new cross section prediction code was needed to predict the cross sections for these new reactions.

2.2 HIVAP

The HIVAP code was produced in the attempt to predict production cross sections for a variety of different reactions. One of the failures of earlier codes like JORPLE and SPIT is their tendency to be useful for only a given region of the Chart of Nuclides, like the heaviest elements. Their approach is based on the black box model of compound nucleus formation and therefore the production cross sections that these codes produced are often off by orders of magnitude in cross section. Further, JORPLE and SPIT are small parameter codes, meaning the only input is the projectile and target atomic number and atomic mass. Inevitably, more parameters would enhance a code's ability to accurately predict production cross sections. As the search for heavier elements moved to the cold fusion reaction mechanism, a different production cross section code was needed.

The HIVAP code is a modification of an earlier code that examined the deexcitation of highly excited nuclei [Gro1967]. This code, named GROGI, is based on
looking at nuclei that have large excitation energies and large angular momentum values.

It incorporates the statistical de-excitation of the excited compound nucleus through
neutron, gamma-ray and charged particle emission. HIVAP is an improvement of the
GROGI code through the incorporation of fission into the de-excitation step as well as the
incorporation of new insights into level density calculations, interaction barriers, groundstate masses and shell-effects and fission barriers. One of the advantages of the HIVAP
code is the use of multiple independent sources for the determination of the many
parameters involved in the calculation of the production cross sections [Ver1984].

The HIVAP code is a modular code that can be used in a multitude of situations. Because of this, a detailed list of the parameters used must be kept in order to distinguish the results of one parameter set from another. The multiple parameters used in the HIVAP code can be adapted to a specific section of the Chart of Nuclides to make cross section predictions for a small region more accurate than the earlier more global codes JORPLE and SPIT.

An overview of the calculations in the HIVAP code used in this dissertation is as follows. Most of the following conditions for HIVAP were set according to calculations done by Reisdorf and Schädel, fitting HIVAP results to actinide target based experimental data [Rei1992]. Compound nucleus formation is considered as separate from the de-excitation step. Fusion occurs when the projectile-target system passes the interaction barrier which is calculated using the Bass interaction potential [Bas1977]. Below the barrier, a WKB (Wentzel-Kramer-Brillouin) approximation is made to estimate barrier penetration. The fusion barrier in HIVAP is considered to be fluctuating using a Gaussian parameter with a user-defined standard deviation. Corrections in the entrance channel are also made for extra push and extra-extra push effects [Swi1982]. The de-excitation of the excited compound nuclei is calculated using the following sources: liquid drop masses [Mye1966], level density calculations [Rei1981], level density ratios [Tök1981], and fission barriers [Coh1974]. The standard set of parameters used from this description of the HIVAP code will be referred to as the Reisdorf and Schädel parameters and are given in Table 2.1.

Table 2.1: Reisdorf and Schädel Parameter set for the HIVAP code [Rei1992].

Variable	Description	Value
LEVELPAR	Scale parameter for the level density	1.153
AF / AN	Level density ratio parameter value	1
BARFAC	Scale parameter for the fission barrier	1
EDAMP	Shell effect damping energy (MeV)	18.0
DELT	Nuclear pairing correction energy (MeV)	11.0
V0	Initial value of the nuclear potential (MeV)	70.0
R0	Nuclear radius parameter (fm)	1.12
D	Fuzziness in the nuclear radius parameter (fm)	0.75
Q2	Nuclear quadrupole moment (fm²)	1050
CRED	Scale parameter for the interaction barrier	1.0
SIGR0	Fluctuation of the interaction barrier (% of R0)	3.0
CUTOFF	Integration limits in (SIGR0) for barrier fluctuations	5.0
XTH	Extra push theory threshold fissility parameter	0.7
APUSH	Slope coefficient from extra push theory	18.0
FPUSH	Angular momentum coefficient from extra push theory	0.75

Additional parameters from the Reisdorf and Schädel parameter set can be seen in the dissertation of Dressler [Dre1999]. Another invaluable source of information regarding the description of the calculation loop and the various parameters and their associated meanings is a guidebook written by Reisdorf [Rei1990].

With the numerous independent variables that can be changed and altered to suit the needs of the experiment, the HIVAP code is definitely not a general global code for predicting production cross sections. The HIVAP code is much more effective at predicting cross sections when the individual parameters are tailored to a specific type of reaction or particular region of the Chart of Nuclides. The aforementioned Reisdorf and Schädel parameters will be used by HIVAP in this dissertation to compare with the experimentally obtained production cross sections. By using codes such as HIVAP to accurately predict production cross sections, information regarding the physical properties of the transactinide nuclei will be obtained.

3 Experimental procedures

As hot- and cold fusion reactions are both used to produce isotopes of the heaviest elements, so are there different experimental procedures used to produce, separate and detect these heavy element isotopes. Three different experimental procedures were used to produce the heavy element isotopes studied for this dissertation.

For each of these individual experimental procedures there are five basic components: the accelerator, the target chamber and targets, the transportation/separation/collection systems, the detector system and the data acquisition system.

The first of the five basic components is the accelerator. The 88-Inch Cyclotron at Lawrence Berkeley National Lab provided all of the heavy ion beams used in the three experimental procedures. The 88-Inch Cyclotron is a sector focused cyclotron that is capable of providing intense beams of a multitude of ions from protons through uranium. The intense beams are produced in advanced electron-cyclotron resonance (ECR) ion sources created and developed at Lawrence Berkeley National Lab. The beams produced by the cyclotron are controlled, directed and maintained by the experienced staff of the 88-Inch Cyclotron.

Two of the experimental procedures involved stationary targets. The compound nucleus recoils from the projectile-target reactions for the first of these experimental procedures were transported to our rotating wheel system known as the Merry-Go-Round (MG) [Hff1980]. The MG system is important in the study of the heavier elements as the length of time between production and detection is extremely short, however, without any chemical separation, the presence of any interfering activity would mask the activity of interest making positive identification difficult. These negatives are balanced by the

positive aspect that the techniques can be used in reactions where highly active actinide targets are required. Only one experiment was performed using this procedure: $^{248}\text{Cm}(^{15}\text{N},\text{xn})^{263-x}\text{Lr}$.

The compound nucleus recoils from the second stationary target experimental procedure were collected on gold catcher foils located directly behind the standing target. The success of the catcher foil technique lies in the ability to effectively chemically separate the activity of interest from the gold foil and the rest of the reaction products. The duration of the chemistry separation is also extremely important, as longer chemical separations lead to the inability to study short lived isotopes. Only one experiment was performed using this procedure: ²³⁸U(¹⁸O,4n)²⁵²Fm.

If non-active or only slightly active targets can be used, a faster and more efficient experimental procedure can be used, the Berkeley Gas-filled Separator (BGS) [Nin1999]. Six separate experiments (238 U(22 Ne,xn) $^{260-x}$ No, 208 Pb(48 Ca,xn) $^{256-x}$ No, 208 Pb(50 Ti,xn) $^{258-x}$ Rf, 208 Pb(51 V,2n) 257 Db, 209 Bi(50 Ti,xn) $^{259-x}$ Db, and 209 Bi(51 V,2n) 258 Sg) were performed using this third experimental procedure involving a rotating target wheel system and the BGS physical separation apparatus.

3.1 MG system

The ²⁴⁸Cm(¹⁵N,3n)²⁶⁰Lr reaction was studied using the Merry-Go-Round (MG) rotating wheel collection and detection apparatus. The target system used a 2.63 mg/cm² beryllium vacuum window, 0.3 mg/cm² nitrogen as a cooling gas, and a 2.58 mg/cm² beryllium target backing. The 0.873 mg/cm² curium target (96% ²⁴⁸Cm, 4% ²⁴⁶Cm) contained approximately 0.84 mg/cm² ²⁴⁸Cm as the oxide ²⁴⁸Cm₂O₃. Because of the short range of the ¹⁵N projectiles in curium, the effective thickness of the curium target was only 0.368 mg/cm². The uncertainty in the target thickness was estimated to be 0.010 mg/cm². The ²⁴⁸Cm had been electroplated [Aum1974, Mül1975] onto the beryllium target backing in a circle 6-mm in diameter. After the compound nucleus reaction, the recoiling products were thermalized in 1 atm of helium gas inside the target chamber. There they were attached to KCl aerosols in a helium gas-jet and transported via a 7meter 1.4-mm i.d. capillary at a flow rate of 1.8 STP liters/min at a pressure of 5.0 PSIG (pounds per square inch gauge) to the MG collection site. A schematic illustration of the target chamber can be seen in Figure 3.1. The activity-laden aerosols were deposited on polypropylene foils $(40-60 \mu g/cm^2 thick)$ held in eighty collection positions located around the periphery of the 51.0-cm diameter fiberglass wheel. The polypropylene foils containing the activity were stepped between six opposing pairs of PIPS (Passivated Implanted Planar Silicon) detectors. Figure 3.2 shows a schematic diagram of the MG collection wheel inside a vacuum chamber.

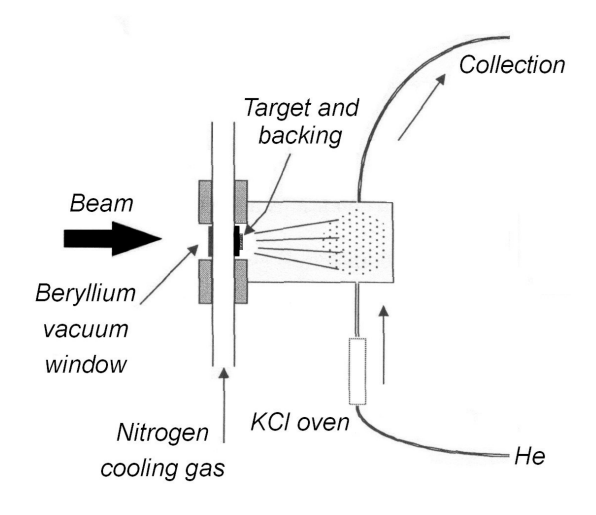


Figure 3.1: Target chamber used in the MG experiment. Recoiling products are thermalized and attached to KCl aerosols in a helium gas-jet for transport to the collection site.

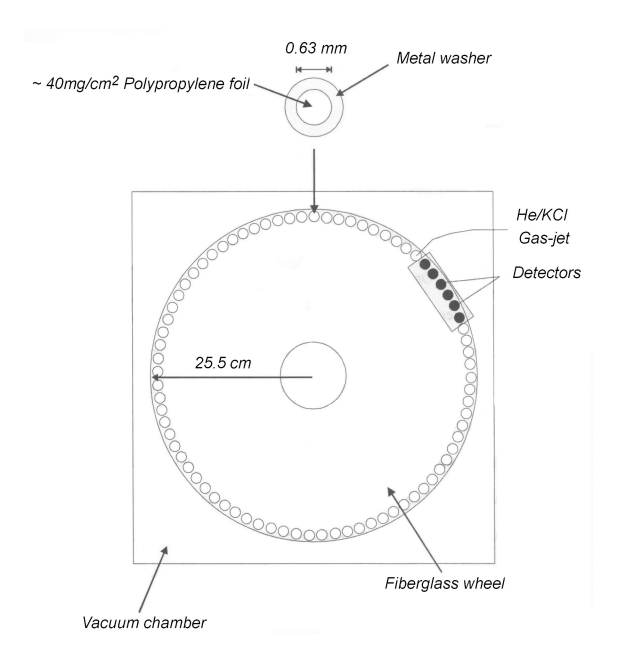


Figure 3.2: View of the Merry-Go-Round rotating wheel collection and detection system. The activity laden KCl aerosols are deposited on one of eighty polypropylene foils located on the periphery of the wheel that is rotated clockwise through the six pairs of detectors, above and below the wheel.

Detectors both below and above the polypropylene foil allowed for coincidence measurements and better efficiency for detecting α -particles. The experimental resolution (FWHM) of the first top detectors was determined to be 45 ± 5 keV by examining the 213 Fr (E_{α} = 6.7750(17) MeV) peak in the first top detector. The experimental resolution for the remaining top detectors was then assumed to be 45 ± 5 keV as well. Experimental resolution for the bottom detectors was approximately 100 keV due to energy degradation in the polypropylene foil [Sha2000]. Transport time from the target to the first polypropylene foil was determined to be 1.0 ± 0.3 s [Lan1998]. A gas-jet and deposition efficiency of $50 \pm 20\%$ was determined by comparing on-line and off-line measurements of 254 Fm (E $_{\alpha}$ = 7.192(2) MeV, $t_{\frac{1}{2}}$ = 3.240 \pm 0.002 h) produced via transfer reactions with the ²⁴⁸Cm target. Measurements were also made of the 4n-out product ^{259}Lr (E $_{\alpha}$ = 8.450(20) MeV, $t_{\frac{1}{2}}$ = 6.3 $^{+0.5}$ $_{-0.4}$ s) to check the accuracy of the gas-jet and deposition efficiency measurement. With a larger cross section and a shorter halflife, several short measurements were made to observe 259 Lr and confirm $50 \pm 20\%$ as the gas-jet and deposition efficiency. The specific details of this measurement are located in Section 4.1. The time between successive wheel movements, or stepping time, was chosen to be two minutes. This allowed for a total detection time of twelve minutes, or four half-lives of 260 Lr (E $_{\alpha}$ = 8.030(20) MeV, $t_{1/2}$ = 180 \pm 30 s). The collection of data was suspended for the first two seconds for the first detector pair to eliminate the detection of the short-lived activities. The uncertainties in the stepping time, collection time, and delayed start counting time were all assumed to 1 millisecond. Given a sourceto-detector distance of two millimeters and an active detector area of 100 mm², a

geometric detector efficiency for a point source of 33% was calculated. The uncertainty in the detector efficiency was assumed to be 2%. New fiberglass wheels with clean polypropylene foils were used every two to three hours to eliminate build up of long-lived activities. The PIPS detectors were calibrated using activity from a 228 Th source. The 228 Th source produces 212 Bi ($E_{\alpha}=6.062(1)$ MeV) and 212 Po ($E_{\alpha}=8.78437(7)$ MeV) activity. The decay data from the 12 detectors, 6 top and 6 bottom, were recorded by the CHAOS [Rat1991] data acquisition software in a list mode that included the detector number, channel, and time for each alpha particle detected. Detection with the six pairs of silicon detectors allowed construction of decay curves for half-life analysis. The CHAOS program generated histogram files used for α -spectrum analysis, α - α correlation analysis and decay-curve analysis.

The production cross sections for the MG experiment were calculated with Equation 3.1,

$$\sigma = \frac{R}{N_t I} \tag{3.1}$$

where R equals the rate that the species of interest is produced, N_t equals the number of target nuclei per unit area, and I equals the rate of the incoming beam particles. This equation was corrected for the MG experiment by accounting for the various decay and growth times (t_{trans} , t_{start} , t_{col} , and t_{count}), efficiencies (Eff_{gj} and Eff_{det}) and branching ratios (BR):

$$\sigma = \frac{N_{events} \lambda}{N_t I_a \left(e^{-\lambda t_{trans}}\right) \left(e^{-\lambda t_{start}}\right) \left(1 - e^{-\lambda t_{col}}\right) \left(1 - e^{-\lambda t_{col}}\right) Eff_{gj} Eff_{det} N_{expt} BR}$$
(3.2)

where N_{events} is the number of events detected in N_{expt} number of similar experiments. λ is the decay constant for the particular isotope of interest.

3.2 Catcher foil experiment

The $^{238}\text{U} + ^{18}\text{O}$ reaction was studied using the catcher foil technique. The target system used a 1.8 mg/cm² HAVAR vacuum window, 0.3 mg/cm² nitrogen as a cooling gas, 2.35 mg/cm² beryllium used as a target backing and a 0.540 mg/cm² ^{238}U target. Correcting for the range of the oxygen ions in uranium, the effective thickness of the uranium target was 0.318 ± 0.010 mg/cm².

The ^{nat}U₃O₈ target was electrodeposited in a manner similar to the curium target used in the MG experiment in Section 3.1. Natural uranium (99.28% ²³⁸U and 0.71% ²³⁵U) as uranylnitrate hexahydrate (UO₂(NO₃)₂•6H₂O) was dissolved in concentrated HCl. The uranium was purified by passing the HCl solution through a Dowex AG1-X8 (200-400 mesh) anion exchange column to sorb the uranium, allowing most impurities to pass through the column. The uranium was eluted from the column with 0.1M HCl and collected. A 10-µl aliquot of the purified uranium solution was evaporated on a platinum disk and counted using alpha spectroscopy to determine the amount of uranium activity per volume of solution. 2077 counts of 238 U ($t_{1/2} = (4.468 \pm 0.003) \times 10^9$ y) were detected in 10 minutes at a detector efficiency of 20% resulting in 17.3 disintegrations of ²³⁸U per second or 1.44 mg of ²³⁸U per 10 µl aliquot. A uranium isopropanol stock solution was made for use in electroplating [Sha2000]. A solution was prepared in which 20 µl of uranium isopropanol solution contained 28.8 µg of ²³⁸U which equaled a thickness of about 0.100 mg/cm² with a target diameter of 6 mm. The 2.35 mg/cm² beryllium target backing was placed in the bottom of the electroplating cell. 20 µl of uranium isopropanol stock solution and 1 ml of isopropanol were placed in the cell. The UO_2^{2+} ions in the

solution were deposited on the beryllium cathode when a 600 V voltage was applied to the anode. The electrodeposition was continued for 45 minutes, the isopropanol solution was removed and the resulting uranium on the beryllium target backing were baked in an oven at 400-500°C for 30 minutes to convert the uranium to the oxide (U_3O_8) form. The thickness of each of the target layers was measured via alpha spectroscopy. The target was placed in a small vacuum chamber under a silicon surface barrier detector at $20 \pm 2\%$ efficiency. After counting the target overnight, the target thickness was determined from the measured 238 U α -activity. A total of seven layers were added to the target by repeating the above process until the target reached a thickness of 0.540 mg/cm². The effective target thickness was 0.318 ± 0.010 mg/cm².

The compound nucleus recoils from this reaction were collected in a 4.69 mg/cm² thick gold catcher foil located five millimeters behind the target in the target chamber. At this thickness, the gold catcher foil collected everything recoiling from the target. The uranium target was irradiated for about 8 hours at each of the energies. The gold foils were carefully removed from the target chamber and then underwent chemical separation. An illustration of the target chamber used in this experiment is shown in Figure 3.3.

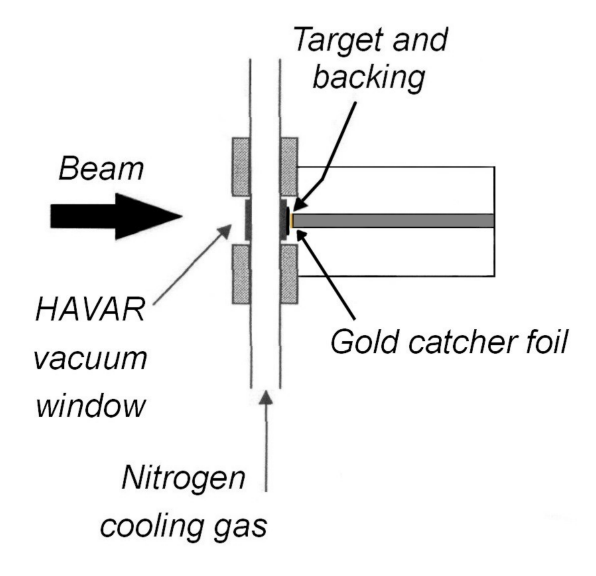


Figure 3.3: Target chamber used in catcher foil experiment. System for the transportation of activity-laden KCl aerosols from Figure 3.1 has been removed and replaced with a gold catcher foil placed directly behind the target.

The first step in the chemical separation was the separation of the trivalent actinides from the gold foil and other reaction products. A flow chart for the chemical separation procedure can be found in Figure 3.4. The highly active gold foil was placed in a test tube containing ^{241}Am (E_{\alpha} = 5.48556(12) MeV, $t_{1/2}$ = 432.2 \pm 0.7 y) tracer and dissolved in 3 drops of concentrated HCl and one drop of concentrated HNO₃. After dissolution, the solution of trivalent actinides was transferred to a Dowex AG1-X8 (200-400 mesh) anion exchange column. The gold and unwanted reaction products were sorbed on the column in concentrated HCl while the trivalent activities passed through. The trivalent actinide solution was dried, taken up in 0.5 M HCl, and transferred to a pretreated Dowex 50W-X4 (200 mesh) cation exchange column. The trivalent fermium activity was separated from the other trivalent activities by elution with 0.5M ammonium alpha-hydroxyisobutyrate (α-HIB) solution at a pH of 3.38 [Cho1956a, Cho1956b, Smi1956]. The fermium fraction eluted from the column in the first 20 drops (4 free column volumes) ahead of the lighter trivalent actinides. Individual drops from the cation column were collected two at a time onto platinum disks. The appropriate platinum disks corresponding to the fermium fraction were transferred to one platinum disk, evaporated to dryness and flamed to fix the activity to the platinum. The ²⁴¹Am tracer was then eluted from the column using 0.5M α -HIB at a pH of 3.72, collected and counted using alpha spectroscopy to determine the chemical separation efficiency. The chemical efficiency was determined to be $90 \pm 10\%$ for the combined anion column and cation column separation. The chemical separation efficiency for the anion column chemistry alone was $95 \pm 5\%$.

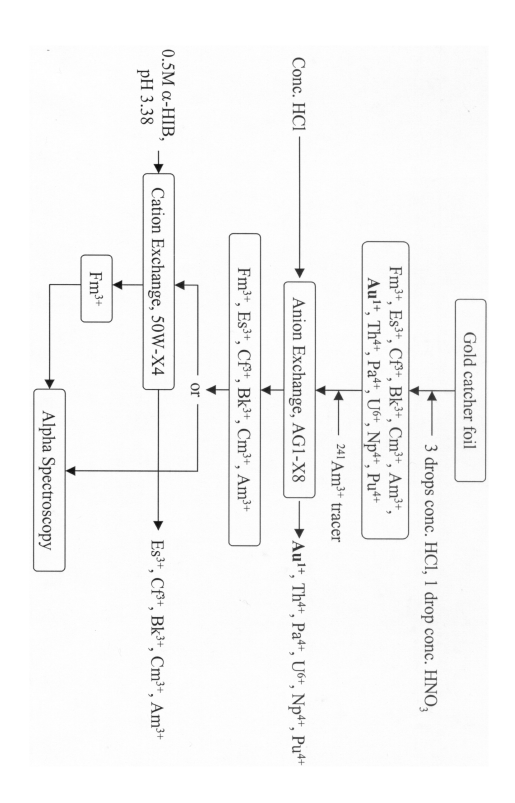


Figure 3.4: Flow chart for the chemical separation of ²⁵²Fm from the gold catcher foil and other unwanted reaction products.

The platinum disk with the fermium fraction was then counted by alpha spectroscopy to observe the decay of fermium. The detector efficiency was $30 \pm 2\%$. Alpha spectra were collected continuously every 3 hours for the first 3 days, then continuously every day for the next seven days. Finally, spectra were obtained continuously every 3 days for the final 20 days. These times correspond to the half-lives of 252 Fm ($t_{1/2} = 25.39 \pm 0.05 \text{ h}$) and 253 Fm ($t_{1/2} = 3.00 \pm 0.12$ d), the 4n- and 3n- exit channel products. The detectors used in these experiments were calibrated using a three peak standard containing ²⁴¹Am, ²⁴⁴Cm $(E_{\alpha} = 5.80482(5) \text{ MeV})$, and ^{252}Cf $(E_{\alpha} = 6.11824(4) \text{ MeV})$. Checks of the calibration were performed during the experiment using activity produced in the experiment. From these individual spectra, decay curves were obtained to determine the initial activity and half-life of ²⁵²Fm. Histogram files were obtained for both experiments using a simple multi-channel analyzer computer card and software from the detector manufacturer. The anion column separation from the gold foil and other reaction products was done for all of the experiments studied. The cation column separation was only performed for the 94.9 MeV experiment.

Cross sections for the catcher foil experiment were measured using the initial activity (A_b) obtained from decay curve fitting and Equation 3.3 which was based on Equation 3.1 corrected for growth and delay times (t_{irr} and t_{start}), efficiencies (Eff_{sep} and Eff_{det}), and branching ratio (BR).

$$\sigma = \frac{A_b}{N_t I_a \left(1 - e^{-\lambda t_{irr}}\right) \left(e^{-\lambda t_{start}}\right) Eff_{sep} Eff_{det} BR}$$
(3.3)

3.3 Berkeley Gas-filled Separator experiments

Six experiments were performed with the BGS: ²⁰⁸Pb(⁵¹V,2n)²⁵⁷Db, ²⁰⁸Pb(⁴⁸Ca,xn)^{256-x}No, ²⁰⁹Bi(⁵⁰Ti,xn)^{258-x}Db, ²⁰⁸Pb(⁵⁰Ti,xn)^{258-x}Rf, ²³⁸U(²²Ne,xn)^{260-x}No, and ²⁰⁹Bi(⁵¹V,2n)²⁵⁸Sg. A general description of the BGS experimental procedure follows [Nin1999]. A schematic diagram of the BGS can be found in Figure 3.5.

Accelerated projectiles passed through a carbon vacuum window that separated the vacuum of the cyclotron and the beam line from the BGS. The beam then passed through a centimeter of helium before hitting the target backing and then the target. The targets in BGS experiments were located on a rotating wheel. Rotating targets were used in these reactions so beam currents larger than those used for normal stationary target reactions could be used. Increased beam currents lead to larger production rates for the activities of interest. By rotating the targets, the heat associated with an intense beam can be spread over several targets instead of one stationary target. Table 3.1 contains the carbon vacuum window thicknesses, carbon target backing thicknesses, target thicknesses, and He pressures in the BGS for the six experiments listed in the order they were performed.

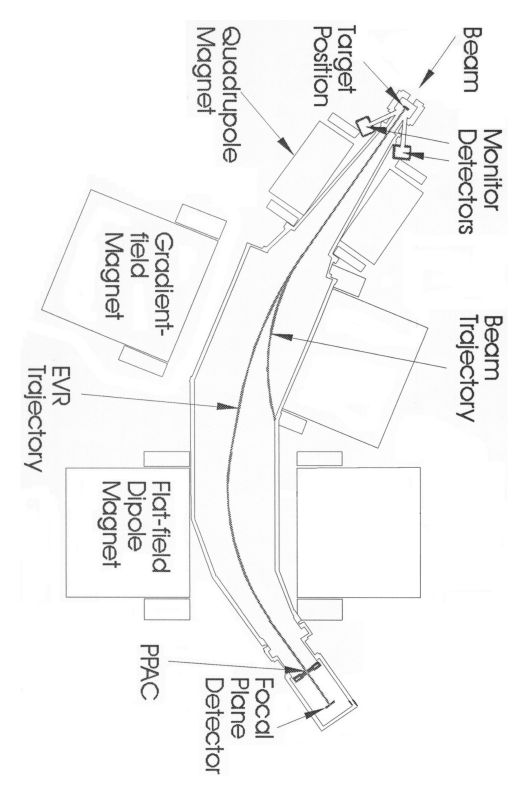


Figure 3.5: Schematic of the BGS showing the target chamber, quadrupole magnet, two dipole magnets and detection chamber. Evaporation residues travel a distance of 460 cm between the target and the PPAC detector.

Table 3.1: BGS reaction specifics. For each reaction, the thickness of the carbon window, carbon backing and target, as well as helium pressure in the BGS is listed. The uncertainty in the target thicknesses were assumed to be $10~\mu g/cm^2$.

	Thickness (μg/cm ²)			
Reaction	Carbon Window	Carbon Backing	Target	He Pressure (Torr)
²⁰⁸ Pb(⁵¹ V,2n) ²⁵⁷ Db	50	35	500	1.000
²⁰⁸ Pb(⁴⁸ Ca,xn) ^{256-x} No	50	35	453	0.742
²⁰⁹ Bi(⁵⁰ Ti,1n) ²⁵⁸ Db	52	40	390	0.785
208 Pb(50 Ti,xn) $^{258-x}$ Rf	50	35	460	1.000
²⁰⁹ Bi(⁵⁰ Ti,xn) ^{259-x} Db	46	29	400	1.000
²⁰⁸ Pb(⁴⁸ Ca,xn) ^{256-x} No	48	35	460	0.787
²³⁸ U(²² Ne,xn) ^{260-x} No	46	46	160.5	0.501
²⁰⁹ Bi(⁵¹ V,2n) ²⁵⁸ Sg	46	26	400	0.712

The ²⁰⁸Pb, ²⁰⁹Bi, and ²³⁸U target wheels were produced either at the Gesellschaft für Schwerionenforschung (GSI) in Darmstadt, Germany or at the Thin Foil Lab at Lawrence Berkeley National Laboratory. Thin carbon target backings were created first and transferred to one of nine target frames. The target frames were banana shaped and measured 105 mm from end to end and 12 mm wide. ²⁰⁸Pb, ²⁰⁹Bi, or ²³⁸U was then evaporated onto the carbon backing producing targets that were uniform in thickness throughout the target wheel.

Beam, transfer products, and compound nucleus evaporation residues (EVRs) from the associated projectile/target reactions traversed the helium gas-filled chamber between the magnets. Rutherford-scattered beam projectiles were used to monitor the amount of beam delivered to the target. Two small p-i-n diode detectors, located 292 ± 1 mm from the target, at an angle $\pm 27.0 \pm 0.1^{\circ}$ from the beam axis, covered by a collimator with a radius of 0.52 ± 0.01 mm, were used to detect the Rutherford scattered beam (see Figure 3.6). The integrated spectra from these detectors was used with the known equations for Rutherford scattering (Equations 3.10 - 3.12) to determine the total beam dose. The scattered beam, transfer products, and the EVRs traveled 460 centimeters through a vertically focusing quadrupole magnet, a gradient field dipole magnet, and a flat field dipole magnet. The quadrupole magnet focused all of the beam, transfers, and EVRs into a rough horizontal plane. The gradient dipole magnet continued the horizontal focusing and began the separation of the beam, transfers and EVRs according to their average charge state and momentum. The flat field magnet then completed the separation.

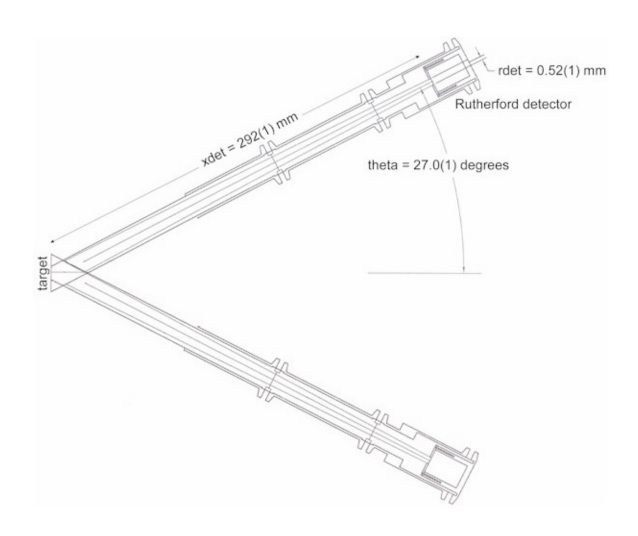


Figure 3.6: Drawing of the Rutherford detector setup showing the target and Rutherford detector positions, distance between the target and Rutherford detector (xdet), the angle between the Rutherford detectors and the beam axis (theta, θ), and the radius of the collimator that covers the Rutherford detectors (rdet). The uncertainties in the last digit of xdet, rdet and theta are listed in parenthesis.

Separation in a gas-filled magnetic separator is dependent on the magnetic field, the velocity and mass of the particle, and its average charge state. Charged particles in a magnetic field feel a force equal to the product of the particle's charge (q), the particle's velocity (v), the magnetic field (B) present, and the sine of the angle (θ) between the velocity and magnetic field vectors.

$$F_B = qvB\sin(\theta) \tag{3.4}$$

This force is equal to the centrifugal force felt by a particle moving in a circle,

$$F_c = \frac{mv^2}{r} \tag{3.5}$$

where m is the mass of the particle and r is the radius of the circle the particle is moving around. Setting these two equations equal to each other and solving for the magnetic rigidity, which is the product of the magnetic field and the radius of curvature for the particle in the magnetic field, gives Equation 3.6.

$$B\rho = \frac{mv}{q} \tag{3.6}$$

The average charge state of the particle moving through a dilute gas can be approximated using the following equation [Bet1972],

$$\overline{q} = Z \left\{ 1 - C_1 \exp \left[-C_2 \left(\frac{v}{v_o} \right) Z^{-\frac{2}{3}} \right] \right\}$$
 (3.7)

where Z is the atomic number of the particle and C_1 and C_2 are constants determined from fits to experimental data. In dilute helium gas [Ghi1988], C_1 and C_2 were

determined to be 1.04 and 0.91 respectively. The Bohr velocity (v_o) in Equation 3.7 is equal to,

$$v_o = \frac{e^2}{2\varepsilon h} \tag{3.8}$$

where e is the elementary charge constant, ε is the permittivity of vacuum and h is Planck's constant. The mass and velocity of Equation 3.6 can be found easily from the kinetics of the observed reaction. Therefore, the beam, transfer products and EVRs can be separated based on their differences in magnetic rigidity in the dilute helium gas.

After separation from the transfer products and beam, the EVRs continued to the detection chamber. The total transit time in the BGS between the target and detector is on the order of 1 μs . Once inside the detection chamber, they passed through a parallel plate avalanche counter (PPAC) before striking the focal plane silicon strip detector. The PPAC was used to provide a time of flight signal. This signal was used to discriminate between events that originated in the focal plane detector and those that originated from EVRs implanted in the detector. The 116-mm wide by 58-mm tall silicon strip detector, 300 μ m thick PIPS detector was divided into 32 vertical strips that recorded energy, time and position, through resistive readout from both the top and bottom of each strip, for each event that hit the detector. The focal plane detector was located towards the back end of the detection chamber, centered between the top, bottom, left and right sides of the detection chamber. The focal plane alpha detection efficiency was $50 \pm 2\%$ and spontaneous fission detection efficiency was $100 \pm 2\%$.

Either a microcomputer called the CVC (CAMAC to VSB Computer) running in a CAMAC (Computer Aided Measurement And Control) crate or the microcomputer RIO2 running in a VME (VERSA Module Eurocard) crate along with the data acquisition software called MBS [Ess2000] were used to collect the data into events and then send it either for storage on tape or for online analysis. The CVC was used for several of the experiments and the RIO2 was used for the rest of the experiments. The RIO2 offered faster data collection and transfer rates than the CVC. A Digital Equipment VAX workstation running the VMS operating system and the data analysis software GOOSY [Ess1987] was used to perform online and offline analysis of the data. For each event over 250 pieces of data or words (see Table 3.2) were recorded. GOOSY analysis routines were written in the PL1 programming language to analyze the data event by event. GOOSY analysis codes were also used to analyze the data from tape after the experiment was completed. The analysis routines were written to produce EVR, alpha, fission, time, position, Rutherford, and PPAC detector spectra as well as EVR-alpha, EVR-fission, and alpha-alpha correlation lists. From these analysis programs, detailed information could be gathered about a particular isotope's decay energies, half-life, activity and eventually production cross section. An example of a GOOSY analysis code can be found in Appendix A.

Table 3.2: Event word list for the ²⁰⁹Bi(⁵¹V,2n)²⁵⁸Sg experiment.

Word Parameter	Description
1-2	microsecond scaler chopper
3-4	microsecond scaler since start acquisition
5-6	millisecond scaler since start acquisition
7-8	second scaler since start acquisition
9-10	minute scaler since start acquisition
11-12	microsecond scaler since @start
13-14	millisecond scaler since @start
15-16	second scaler since @start
17-18	minute scaler since @start
19-20	number of beam pulses
21-22	number of beam dumps
23-24	scaled number of beam dumps
25-26	Rutherford east scaler
27-28	Rutherford west scaler
29	user bit 0,1
30	error location
31	error type
32-63	energy low focal plane detector strips 1-32
64-95	energy high focal plane detector strips 1-32
96-127	position low top focal plane detector strips 1-32
128-159	position high top focal plane detector strips 1-32
160-191	position low bottom focal plane detector strips 1-32
192-223	position high bottom focal plane detector strips 1-32
224-239	energy low backward detector strips 1-16
240-255	energy high backward detector strips 1-16
256-259	individual PPAC signals
260	PPAC time to amplitude converter signal
261-267	- empty -
268	energy low sodium iodide detector
269	energy high sodium iodide detector
270	energy Rutherford east detector
271	energy Rutherford west detector
272-279	energy low punch through detector strips 1-8

Production cross sections for the BGS reactions were calculated using a different method than what has been described in Sections 3.1 and 3.2. The BGS technique for calculating the production cross section is based on a ratio of the number of events observed (N_{events}) to the number of Rutherford events ($N_{rutherfords}$) observed, multiplied by the Rutherford scattering cross section ($\sigma_{Rutherford}$):

$$\sigma = \frac{N_{events}}{N_{Rutherfords}} \sigma_{Rutherford}$$
(3.9)

The number of events observed and the number of Rutherford-scattered events observed can be found in the experimental data. $\sigma_{Rutherford}$ involves the calculation of the Rutherford scattering differential cross section and then multiplying by the solid angle subtended by the Rutherford detector. The lab frame Rutherford scattering differential cross section equation is found in Equation 3.10 [Seg1977],

$$\frac{\partial \sigma}{\partial \omega_{lab}} = \left(\frac{e^2 Z_p Z_t}{2Ebeam_{lab}}\right)^2 \sin^{-4}(\theta_{lab}) \frac{\left[\cos(\theta_{lab}) + \left[1 - \left(\frac{A_p}{A_t}\right)^2 \sin^2(\theta_{lab})\right]^{\frac{1}{2}}\right]^2}{\left[1 - \left(\frac{A_p}{A_t}\right)^2 \sin^2(\theta_{lab})\right]^{\frac{1}{2}}} (3.10)$$

where Z_p and Z_t are the atomic numbers of the projectile and target and A_p and A_t are the atomic mass numbers of the projectile and target. The angle and beam energy are in the lab frame and are determined in the experiment. The solid angle can be determined from the following equation,

$$\partial \omega_{lab} = \pi \left(\frac{r \det}{x \det} \right)^2 \tag{3.11}$$

where rdet is the radius of the detector and xdet is the distance from the detector face to the target (see Figure 3.6). The product of Equation 3.10 and 3.11 gives rise to $\sigma_{Rutherford}$.

$$\sigma_{Rutherford} = \frac{\partial \sigma}{\partial \omega_{lab}} \partial \omega_{lab}$$
 (3.12)

Add in efficiencies (Eff_{bgs}, Eff_{det} and Eff_{det2}) and the branching ratio (BR) to arrive at the final result for the BGS cross section calculation method.

$$\sigma = \frac{N_{events}\sigma_{Rutherford}}{N_{rutherfords}Eff_{bgs}Eff_{det}Eff_{det2}BR}$$
(3.13)

4 Experimental results

Table 4.1 contains the half-lives, branching ratios and primary α -decay energies for the isotopes studied in our experiments [Chu1999]. All of the beam energies used in this section are measured in the lab frame. Equation 1.26 with a radius parameter (r_e) of 1.4 fm was used to calculate the Coulomb barriers for the reactions discussed. The Coulomb barriers are all listed in the lab frame. Excitation energies are calculated in the center of mass system. Uncertainties in the center of target beam energies were assumed to be 2 MeV. The uncertainties in the number of counts was determined from the results of MLDS fits, the square root of the number of counts (number of counts > 20), or from the statistics of [Sch1984] (number of counts ≤ 20) assuming a 68% confidence interval. Unless otherwise noted, the errors in the experimental cross sections were a combination of statistical errors in the initial activity or number of counts and the systematic errors of the experiment. In the BGS experiments, the uncertainty in the time of each experiment was assumed to be 5 seconds, and the uncertainty in the number of Rutherford counts was assumed to be the square root in the number of Rutherford counts.

Table 4.1: Summary of the half-lives, branching ratios and primary α -decay energies for the isotopes studied in our experiments [Chu1999]. Approximate uncertainties (\sim) in the branching ratios were assumed to be 5%.

Isotope	Half-life	Branching Ratios	E _α (MeV)
²¹⁷ At	$32.3 \pm 0.4 \text{ ms}$	$\alpha 99.988 \pm 0.004\%$	7.0699(5)
²¹³ Rn	$25.0 \pm 0.2 \text{ ms}$	α 100%	8.088(8)
²¹³ Fr	$34.6 \pm 0.3 \text{ s}$	$\alpha~99.45\pm0.03\%$	6.7750(17)
²¹⁴ Ac	$8.2 \pm 0.2 \text{ s}$	α 89 \pm 3%, sc 11 \pm 3%	7.214(5)
²⁵² Fm	$25.39 \pm 0.05 \text{ h}$	$\alpha~99.9977 \pm 0.0002\%$	7.039(2)
²⁵² No	$2.30 \pm 0.22 \text{ s}$	α 73.1 ± 1.9%, SF 26.9 ± 1.9%	8.415(6)
²⁵³ No	$1.7 \pm 0.3 \text{ min}$	α ~80%, ec ~20%	8.010(20)
²⁵⁴ No	$55 \pm 3 \text{ s}$	$\alpha~90\pm4\%$	8.093(14)
²⁵⁵ No	$3.1 \pm 0.2 \text{ min}$	α 61.4 \pm 2.5%, sc 38.6 \pm 2.5%	8.121(6)
²⁵⁶ No	$2.91 \pm 0.05 \text{ s}$	$\alpha~99.5\pm0.1\%$	8.430(20)
²⁵⁷ No	$25 \pm 2 \text{ s}$	α ~100%	8.220(20)
²⁵³ Lr	$1.3^{+0.6}$ $_{-0.3}$ s	α 98 ± 2%	8.800(20)
²⁵⁹ Lr	$6.3^{+0.5}_{-0.4} \text{ s}$	α 77 ± 2%, SF 23 ± 2%	8.450(20)
²⁶⁰ Lr	$180 \pm 30 \text{ s}$	α 75 \pm 10%, sc 25 \pm 10%	8.030(20)
²⁵⁶ Rf	$6.7 \pm 0.2 \text{ ms}$	SF 98 ⁺² -7%	
²⁵⁷ Rf	$4.7 \pm 0.3 \text{ s}$	α 79.6 ± 2.0%, ϵ c 18 ± 2%	8.774(8), 9.013(8)
258 Rf	$12 \pm 2 \text{ ms}$	SF ~87%	
²⁵⁷ Db	$1.3^{+0.5}$ _{-0.3} s	α 82 \pm 11%, SF 17 \pm 11%	8.970(20)
²⁵⁸ Db	$4.4^{+0.9}_{-0.6} \text{ s}$	α 67 $^{+5}$ -9%, sc 33 $^{+9}$ -5%	9.172(15)
258 Sg	~2.9 ms	SF ~100%	

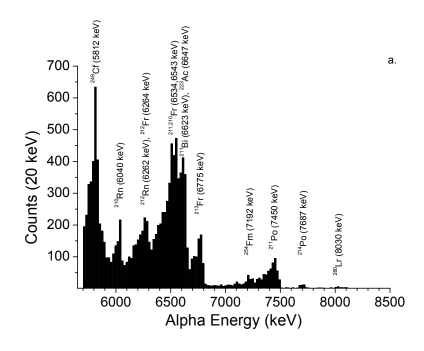
$4.1 ^{248}\text{Cm} + ^{15}\text{N}$

 260 Lr (E_{α} = 8.030(20) MeV, $t_{/2}$ = 180 ± 30 s) and 259 Lr (E_{α} = 8.450(20) MeV, $t_{/2}$ = 6.3 $^{+0.5}$ - $_{0.4}$ s) were produced via the 248 Cm(15 N,xn) $^{263-x}$ Lr reaction, where x = 3 or 4. The MG system described in Section 3.1 was used to produce, transport, collect, and detect the 260 Lr and 259 Lr activity. 15 N⁴⁺ out of the cyclotron energies of 92.0 and 94.5 MeV corresponded to energies of 75.9 and 78.8 MeV in the center of the 0.873 mg/cm² curium target, which corresponded to excitation energies of 35.2 and 37.9 MeV, respectively. The Coulomb barrier for this reaction was calculated to be 83.8 MeV. Fiberglass wheels with clean polypropylene foils were used for each of the energies and isotopes of interest.

4.1.1 ²⁶⁰Lr

Beam doses of $(1.210 \pm 0.004) \times 10^{17}$ (75.9 MeV) and $(1.715 \pm 0.005) \times 10^{17}$ (78.8 MeV) were accumulated for the two experiments, corresponding to average beam currents of 2.688 ± 0.010 and 2.568 ± 0.008 e μ A on target, respectively. There were a total of 240 separate collections at 75.9 MeV and a total of 354 separate collections at 78.8 MeV.

Alpha decays detected by five of the six top detectors for the two experimental energies were summed to produce the spectra in Figures 4.1 and 4.2. The first top detector was not included in these summed α -spectra. Only alpha decays with energies greater than 5.7 MeV were included in these summed spectra to make sure all of the peaks were visible on the same scale. Table 4.2 lists the prominent activities seen in Figures 4.1 and 4.2 with their decay energies, half-lives and possible production routes.



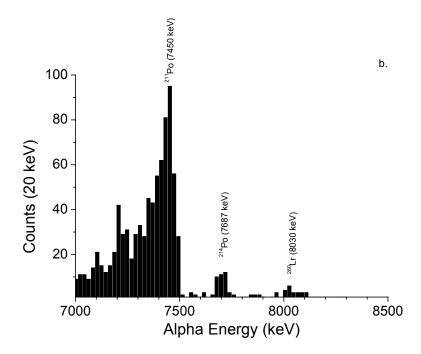
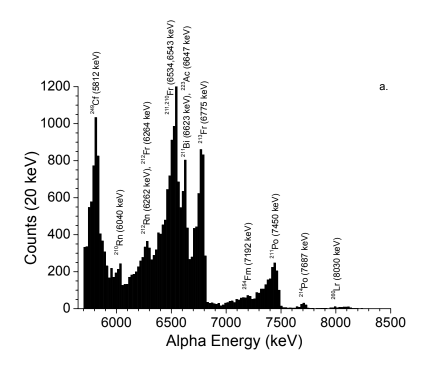


Figure 4.1: Sum spectrum of detectors two top through six top of the MG for the 75.9-MeV 248 Cm + 15 N experiment (a.). Expanded region between 7000 - 8500 keV shown in (b.)



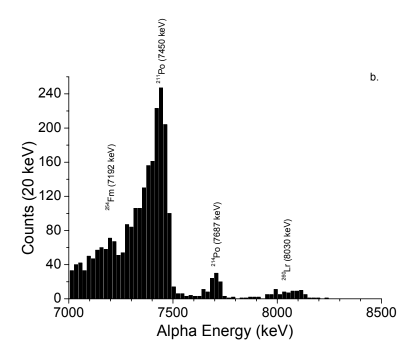


Figure 4.2: Sum spectrum of detectors two top through six top of the MG for the 78.8-MeV 248 Cm + 15 N experiment (a.). Expanded region between 7000 - 8500 keV shown in (b.)

Table 4.2: List of alpha-decay peak assignments for the sum spectra in Figures 4.1 and 4.2. Alpha decay energies and half-lives from [Chu1999].

Isotope	E_{α} (MeV)	Half-life	Possible Production Route
²⁴⁹ Cf	5.8133(10)	$351 \pm 2 \text{ y}$	present naturally, contamination
²¹⁰ Rn	6.040(2)	$2.4 \pm 0.1 \text{ h}$	electron capture decay of ²¹⁰ Fr
²¹² Rn	6.262(3)	23.9 ± 1.2 min	electron capture decay of ²¹² Fr
²¹² Fr	6.2619(21)	$20.0 \pm 0.6 \text{ min}$	$^{206}\text{Pb}(^{15}\text{N},\alpha5\text{n}), ^{207}\text{Pb}(^{15}\text{N},\alpha6\text{n}),$ $^{208}\text{Pb}(^{15}\text{N},\alpha7\text{n})$
²¹¹ Fr	6.534(5)	$3.10 \pm 0.02 \text{ min}$	204 Pb(15 N, α 4n), 206 Pb(15 N, α 6n)
²¹⁰ Fr	6.543(5)	$3.18 \pm 0.06 \text{ min}$	204 Pb(15 N, α 5n), 206 Pb(15 N, α 7n)
²¹¹ Bi	6.6229(6)	$2.14 \pm 0.02 \text{ min}$	208 Pb(15 N,3α), 223 Ac \to α-decay
²²³ Ac	6.6467(10)	$2.10 \pm 0.05 \text{ min}$	$^{208}\text{Pb}(^{15}\text{N},\gamma)$
²¹³ Fr	6.7750(17)	$34.6 \pm 0.3 \text{ s}$	208 Pb(15 N,α6n), 206 Pb(15 N,4n) 217 Ac → 213 Fr + α
²⁵⁴ Fm	7.192(2)	$3.240 \pm 0.002 \text{ h}$	248 Cm(15 N, α 5n) 254 Md \rightarrow 254 Fm, or transfer reaction
²¹¹ Po	7.4503(5)	0.516 ± 0.003 s	208 Pb(α ,n), 207 Pb(α , γ)
²¹⁴ Po	7.68682(7)	$164.3 \pm 2.0 \ \mu s$	present naturally
²⁶⁰ Lr	8.030(20)	$180 \pm 30 \text{ s}$	²⁴⁸ Cm(¹⁵ N,3n)
²¹³ Rn	8.090(8)	$25.0 \pm 0.2 \text{ ms}$	electron capture decay of ²¹³ Fr

Most of the isotopes seen in Table 4.2 are the result of compound nucleus reactions between the nitrogen projectiles and trace amounts of lead impurities in the curium target. Although the amounts of lead in the curium target might be small, the cross sections for these reactions are more than an order of magnitude higher than for the production of ²⁶⁰Lr leading to similar production rates.

The search for ²⁶⁰Lr began by examining the individual singles alpha spectra for the two different energy experiments. In particular, the search was aimed at finding a 180-second activity located around 8.0 MeV. A region of the singles spectrum for each of the two experiments was established around 8.030 MeV and integrated in each of the twelve detectors used in the experiment, six on top and six on bottom. The region of interest chosen for these experiments was 7.95 – 8.15 MeV. With an experimental resolution of 45 keV in the top detectors, this region of interest was sufficient to include the decay of ²⁶⁰Lr. Each top/bottom partner integrated region of interest was summed as they both correspond to the same time window. The decay curve values for the two energy experiments can be found in Table 4.3.

Table 4.3: Decay tables for the two reactions showing the number of counts in the region of interest given for each of the two experiments for each set of detectors. The range of time that the samples were between a given set of detectors is noted next to each detector pair.

	Experiment (Region of Interest)		
	75.9 MeV ¹⁵ N (7.95 - 8.15 MeV)	78.8 MeV ¹⁵ N (7.95 - 8.15 MeV)	
Detector Pair (time, min)	Number of Co	ounts in Region	
1T + 1B (0-2)	89	466	
2T + 2B (2-4)	21	77	
3T + 3B (4-6)	4	15	
4T + 4B (6-8)	6	8	
5T + 5B (8-10)	5	6	
6T + 6B (10-12)	1	4	

This region of interest was extended to 8.15 MeV to include alpha decays from $^{213}Rn~(E_{\alpha}$ = 8.088(8) MeV, $t_{1/2}$ = 25.0 \pm 0.2 ms), the electron capture decay daughter of 213 Fr ($t_{1/2} = 34.6 \pm 0.3$ s). During the experiment, a secular equilibrium is reached between ²¹³Fr and ²¹³Rn. As a result of the secular equilibrium, the ²¹³Rn activity takes on the half-life of ²¹³Fr. Knowing that this region of interest contained the two activities, the decay curves were then analyzed with a two-component fit using the MLDS method [Gre1991]. The MLDS method is a multicomponent decay curve analysis technique that uses the maximum likelihood technique for decay curves made up of time binned events. Believing the ²⁶⁰Lr was indeed present, the half-life of ²⁶⁰Lr was fixed at 180 seconds in the MLDS fit. The initial activity of the ²⁶⁰Lr was allowed to vary. The half-life and activity of the shorter second component were both allowed to vary. The MLDS program found the best half-life and initial activity for each of the components using the input information given. By integrating the resulting best fit decay curve for the MLDS results, the number of counts corresponding to each component was determined. The results of the MLDS fits to the decay curves in Table 4.3 can be seen in Table 4.4.

Table 4.4: Results of MLDS fits [Gre1991] to the decay curves found in Table 4.3. The errors given are also a result of the MLDS program. The half-life of ^{260}Lr was fixed (^{260}Lr $t_{1/2}=180\pm30$ s, $^{213}Rn,^{213}Fr$ $t_{1/2}=34.6\pm0.3$ s).

Beam Energy	Isotope	Half-life	Initial Activity	Number of Counts
75.9 MeV	²⁶⁰ Lr	180 s	11 ⁺³ -2 min ⁻¹	46 +11 -10
	²¹³ Rn	37 ⁺⁶ -5 s	89 ⁺¹⁴ -13 min ⁻¹	80 +13 -12
78.8 MeV	²⁶⁰ Lr	180 s	$19 \pm 4 \text{ min}^{-1}$	78^{+18}_{-16}
	²¹³ Rn	39.8 ^{+2.0} -1.9 s	$520 \pm 30 \text{ min}^{-1}$	500 ± 30

Using the number of counts for 260 Lr obtained from the MLDS fit, a 180 ± 30 s half-life, a $66 \pm 2\%$ α -particle detection efficiency, and an alpha-decay branching ratio of $75 \pm 10\%$, the production cross section could be calculated using Equation 3.2. 260 Lr production cross sections of 2.3 ± 1.2 nb and 2.8 ± 1.4 nb were obtained for the 248 Cm(15 N,3n) 260 Lr reaction at excitation energies of 35.2 MeV and 37.9 MeV, respectively. A previous experiment [Esk1971] that produced 260 Lr in the 248 Cm(15 N,3n) reaction reported a production cross section for 260 Lr of about 2 nb at an excitation energy of 37.2 MeV. Their cross section value is consistent with our experimental cross sections. An excitation function for the 248 Cm(15 N,3n) reaction was calculated using HIVAP with the parameters given in Section 2.2. Our experimental results, the results of Eskola et al., and the results of the HIVAP calculation using the parameters of Reisdorf and Schädel are plotted in Figure 4.3.

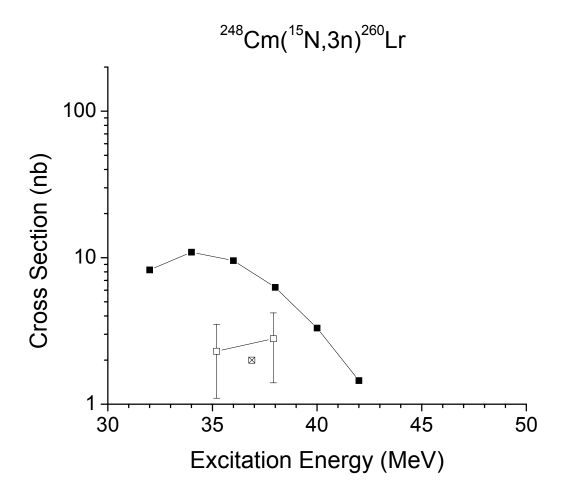


Figure 4.3: Comparison of experimental results and calculated values for the ²⁴⁸Cm(¹⁵N,3n) reaction. Solid symbols (+) represent the HIVAP code using the Reisdorf and Schädel parameters, open symbols (0) represent the experimental results of this thesis, and the open symbols with an (4) through them represent previous experimental results [Esk1971].

$4.1.2^{259}$ Lr

Over the course of one hour, a total of 915 separate collections were performed in the 75.9 MeV experiment, and over the course of several hours a total of 4508 separate collections were performed in the 78.8 MeV experiment. The experimental collection time and stepping time for both experimental energy experiments were altered to correspond to the shorter half-life of 259 Lr compared to 260 Lr. A stepping time of four seconds was used, corresponding to a total time under the detectors of 24 seconds or 4 half-lives of 259 Lr. As with the 260 Lr experiment, there was a two second dead time window for the collection of data under the first detector. The experiment was run for a total of approximately 1 hour at 75.9 MeV and approximately 5 hours at 78.8 MeV, at an average beam current of 2.80 ± 0.03 e μ A and 2.506 ± 0.012 e μ A which corresponded to a beam doses of $(0.1594 \pm 0.0016) \times 10^{17}$ and $(0.706 \pm 0.003) \times 10^{17}$, respectively.

The analysis of this experiment was a little more straight forward than the analysis for the 260 Lr experiment. Very little activity interfered with the 259 Lr activity at 8.450 MeV. The first top and first bottom detector were ignored in response to the large amounts of short-lived activities present in those spectra. Because of its better resolution, the alpha spectrum from the second top detector was integrated around 8.45 MeV and the total number of counts obtained in that integration was used to calculate the production cross section. Because only the alpha-decay recorded from one detector was used, the detector efficiency was only $33 \pm 2\%$. As only the second top detector was used, the time between the end of collection and the beginning of detection and the total collection time was just 4 seconds. The transport time was still 1.0 ± 0.3 s and the gas-jet and deposition efficiency was $50 \pm 20\%$.

The production cross sections at the two energies were calculated assuming an alpha-decay branching ratio of 77 ± 2% and a half-life for ²⁵⁹Lr of 6.3 ^{+0.5}_{-0.4} seconds. Only 8 events were detected for the 75.9 MeV experiment and a total of 122 events were detected for the 78.8 MeV experiment, which leads to production cross sections of 27 ⁺¹⁷₋₁₄ nb and 90 ± 40 nb at excitation energies of 35.2 MeV and 37.9 MeV, respectively. This experiment was also previously measured by Eskola, et al. [Esk1971] and compared favorably with the experimental results obtained in this thesis. The experimental point at 37.9 MeV is higher than what might be expected from the Eskola results, but no cross sections were measured around that energy by Eskola. The experimental cross sections do match fairly well with the HIVAP predictions using the Reisdorf and Schädel parameters. The experimental results for ²⁵⁹Lr, the experimental results of Eskola, et al., and the results of the HIVAP code using the Reisdorf and Schädel parameters are shown in Figure 4.4.

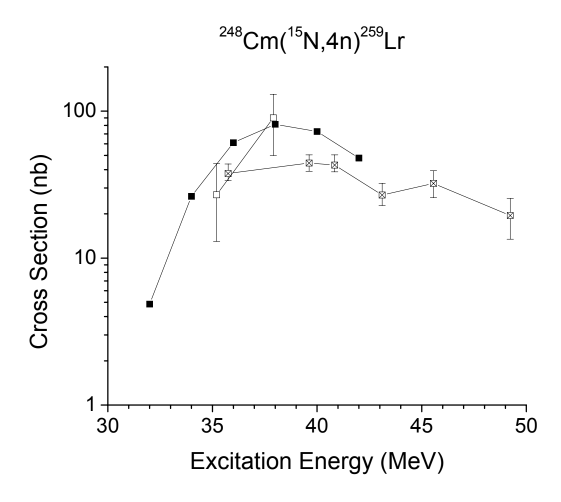


Figure 4.4: Comparison of experimental results and calculated values for the ²⁴⁸Cm(¹⁵N,4n) reaction. Solid symbols (+) represent the HIVAP code using the Reisdorf and Schädel parameters, open symbols (0) represent the experimental results of this thesis, and the open symbols with an (4) through them represent previous experimental results [Esk1971].

$4.2 \quad ^{238}\text{U} + ^{18}\text{O} \rightarrow ^{252}\text{Fm} + 4\text{n}$

²⁵²Fm was produced in the ²³⁸U(¹⁸O,4n) reaction at three different energies of ¹⁸O⁵⁺. The reactions specifics for the ²³⁸U(¹⁸O,4n)²⁵²Fm experiment are located in Table 4.5. The Coulomb barrier for this reaction was 92.3 MeV. After irradiation, the gold foils were then taken and processed chemically to determine the amount of ²⁵²Fm produced.

All of the gold foils were processed chemically according to the procedure described in Section 3.2, separating the ²⁵²Fm from the gold foil and unwanted nontrivalent species. For the 82.4 MeV experiment and the 99.3 MeV experiment, the fermium fraction was then counted without further processing. The chemical efficiency for these single chemistry experiments was $95 \pm 5\%$. The total time between the end of the irradiations and the beginning of counting for these two experiments was 168 ± 5 minutes and 196 ± 5 minutes respectively. For the 94.9 MeV experiment, the fermium fraction was then processed to separate it from the rest of the trivalent species present. This activity was then counted. The chemical separation efficiency for this experiment was $90 \pm 10\%$. The total time between the end of the irradiation and the beginning of counting for the 94.9 MeV experiment was 525 ± 5 minutes. Background spectra for the three detectors used were obtained over a 21 hour period. The detectors for the 82.4 MeV experiment and 99.3 MeV experiment showed no background counts from 6.3 MeV to 10.0 MeV. The detector used for the 94.9 MeV experiment had a few randomly scattered single events between 6.3 MeV and 10.0 MeV. The detectors used had an energy resolution of 85 ± 15 keV. Figures 4.5, 4.6 and 4.7 are cumulative alpha

decay spectra obtained for the three experiments. Table 4.6 lists the isotopes present, their alpha decay energies, and their half-lives. The alpha spectra for the single chemistry experiments show the presence of additional activity resulting from transfer reactions on the uranium target and lead impurities in the target.

Table 4.5: Reaction specifics for the ²³⁸U(¹⁸O,4n)²⁵²Fm experiment. CYC denotes cyclotron energies, COT denotes center of target energies, and E* denotes excitation energies.

E _{CYC} (MeV)	E _{COT} (MeV)	E* (MeV)	Current (eµA)	Dose (×10 ¹⁶)	Time (s)
98.3	82.4	37.7	1.663 ± 0.008	6.00 ± 0.03	28900
109.5	94.9	49.3	1.804 ± 0.008	5.90 ± 0.03	26205
113.5	99.3	53.4	2.397 ± 0.009	9.19 ± 0.03	30700

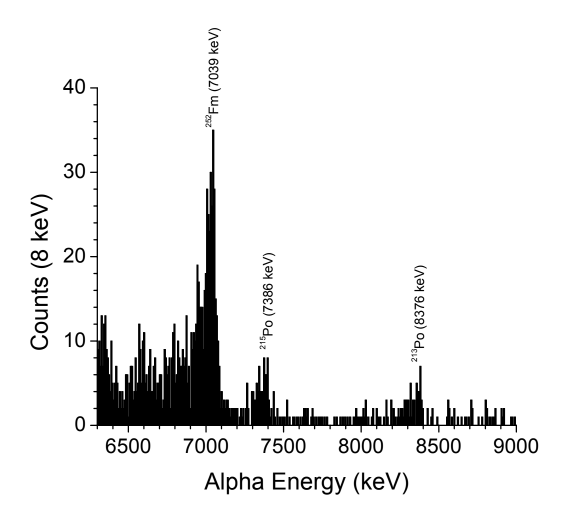


Figure 4.5: Summed alpha spectrum for the 82.4-MeV 238 U + 18 O experiment.

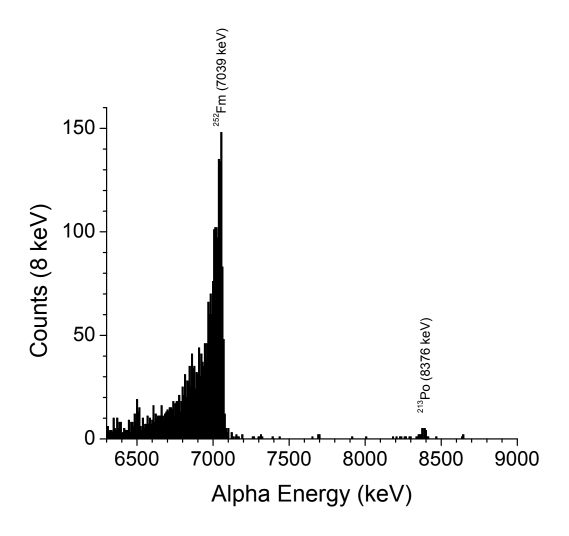


Figure 4.6: Summed alpha spectrum for the 94.9-MeV $^{238}U + ^{18}O$ experiment. Very little activity is present above 7.1 MeV as a result of the α -HIB cation chemistry that was used to separate the fermium from the rest of the trivalent activities and impurities present.

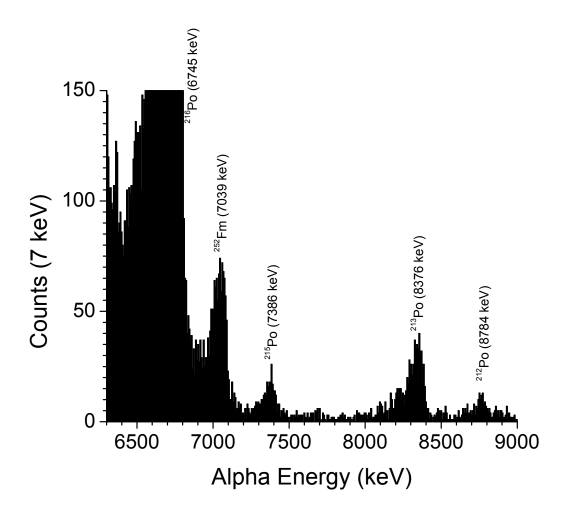


Figure 4.7: Summed alpha spectrum for the $99.3\text{-MeV}^{238}\text{U} + {}^{18}\text{O}$ experiment.

Table 4.6 List of alpha-decay peak assignments for the spectra in Figures 4.5, 4.6 and 4.7. The polonium isotopes result from the decay of long-lived activities produced in transfer reactions with the uranium target and lead impurities in the uranium target. Alpha decay energies and half-lives from [Chu1999].

Isotope	E_{α} (MeV)	Half-life
²¹⁶ Po	6.7783(5)	0.145 ± 0.002 s
²⁵² Fm	7.039(2)	$25.39 \pm 0.05 \text{ h}$
²¹⁵ Po	7.3862(8)	1.781± 0.004 ms
²¹³ Po	8.3759(25)	$4.2\pm0.8~\mu s$
²¹² Po	8.78437(7)	$0.299 \pm 0.002 \; \mu s$

As described in Section 3.2, the samples were counted continuously over various time periods to obtain decay curves. The integration of the same region of interest over various time periods produced a decay curve for each energy. The initial activities of the isotopes present in the region of interest were determined from MLDS fits to the decay data. In the search for 252 Fm ($E_{\alpha} = 7.039(2)$ MeV, $t_{V_3} = 25.39 \pm 0.05$ h), the particular region of interest was approximately between 6.9 and 7.1 MeV. The large search region of interest was due to the poor energy resolution in the detectors (85 ± 15 keV) and alpha particle energy loss in the sample. A time window of 24 hours was chosen to look for the 25.39-hour decay of 252 Fm. The results of the integrations of the specific regions of interest for the three different energy experiments are listed in Table 4.6. Because of the differing times that the experiments occurred, not all of the time bins are exactly 24 hours. This factor is considered in MLDS when determining the half-life and initial activity of the 252 Fm for each experiment.

A two component fit was used to solve for the decay of 252 Fm. The first component would be the 25.39-hour 252 Fm and the second component would be a long-lived component due to the decay of 217 At ($E_{\alpha} = 7.0669(15)$ MeV, $t_{1/2} = 32.3 \pm 0.4$ ms). 217 At is present from the decay of the long-lived 229 Th which is present naturally and reaches an equilibrium with its longer lived parent and grandparents, giving rise to a much longer half-life than 252 Fm. The results of the MLDS fits to the decay curves can be found in Table 4.7. The half-lives and initial activities are the result of the MLDS program. The total number of counts is determined by integrating the MLDS result decay curve over the given counting interval which was 9.53 days (82.4 MeV), 8.75 days (94.9 MeV), and 8.50 days (99.3 MeV).

Table 4.7: Decay tables for the three reactions showing the number of counts in the region of interest given for each of the three experiments. The time window in which the integration of the alpha spectra were integrated is also listed.

Experiment (Region of Interest in MeV)					
82.4 MeV (6.906 – 7.086)		94.9 MeV (6.908– 7.100)		99.3 MeV (6.902 – 7.097)	
Time Window	Number of Counts	Time Window	Number of Counts	Time Window	Number of Counts
0-1 d	163	0-1 d	811	0-1 d	381
1-2 d	90	1-2 d	372	1-1.75 d	177
2-2.83 d	41	2-3 d	196	1.75-2.75 d	101
2.83-3.83 d	22	3-4 d	99	2.75-3.75 d	58
3.83-4.83 d	17	4-5 d	60	3.75-4.75 d	46
4.83-5.83 d	16	5-6 d	36	4.75-5.75 d	23
5.83-6.83 d	3	6-7 d	15	5.75-6.75 d	17
6.83-7.83 d	5	7-8 d	11	6.75-7.75 d	17
7.83-8.83 d	0	8-8.75 d	1	7.75-8.50 d	14
8.83-9.53 d	2			1	

Table 4.8: Results of MLDS fits to the decay curves found in Table 4.6. The errors given are also a result of the MLDS program (252 Fm $t_{1/2} = 25.39 \pm 0.05$ hr). Both the half-life and initial activities were allowed to fluctuate when performing these fits. Fixing the half-life of 252 Fm resulted in differences in the initial activity of only 3%.

Beam Energy	Isotope	Half-life	Initial Activity	Number of Counts
82.4 MeV	²⁵² Fm	27.2 ^{+1.4} _{-1.3} hr	215 ⁺¹⁴ -13 d ⁻¹	350 ± 20
94.9 MeV	²⁵² Fm	24.6 ^{+0.6} -0.5 hr	$1080 \pm 30 \ d^{-1}$	1590 ± 50
99.3 MeV	²⁵² Fm	$22.1 \pm 0.9 \text{ hr}$	$530 \pm 30 \text{ d}^{-1}$	710 ± 30

The results of the MLDS fits to the experimental decay curves show half-lives that are in good agreement with the half-life of 25.39 ± 0.05 hours. Assuming a $99.9977 \pm 0.0002\%$ alpha decay branch, the initial activities from Table 4.7, and the half-life of 252 Fm of 25.39 ± 0.005 hours, production cross sections for the 238 U(18 O,4n) 252 Fm reaction were calculated to be 29 ± 3 nb, 180 ± 20 nb, and 47 ± 5 nb at excitation energies of 37.7 MeV, 49.3 MeV and 53.4 MeV, respectively.

This experiment has been performed experimentally once before. Donets et al., reported cross sections of fermium isotopes produced in the same reaction at energies from 80-135 MeV [Don1966]. The experimental cross sections from this thesis are only comparable at the highest energies for the 4n-reaction and are within a factor of two of the results of Donets. The experimental energies from Donets are reported only as energies of the projectiles in the lab frame. These energies were used to calculate the excitation energies for comparison with the experimental results of this thesis and HIVAP. Donets reported a uranium target thickness of approximately 1 mg/cm². This would lead to energy differences of up to 1 MeV in excitation energy at the center of the target. The Reisdorf and Schädel parameters were also used in the HIVAP code to predict the excitation function for the 4n-exit channel reaction. The experimental results match with the HIVAP predictions at the two highest energies, but is off considerably at the lowest energy. One possible explanation for the difference in the cross sections could be that the chemistry performed at the lowest energy was incomplete. A correctly performed chemistry would have displayed a larger initial activity and therefore cross section. The experimental results, the results of Donets and the HIVAP predictions are located in Figure 4.8.

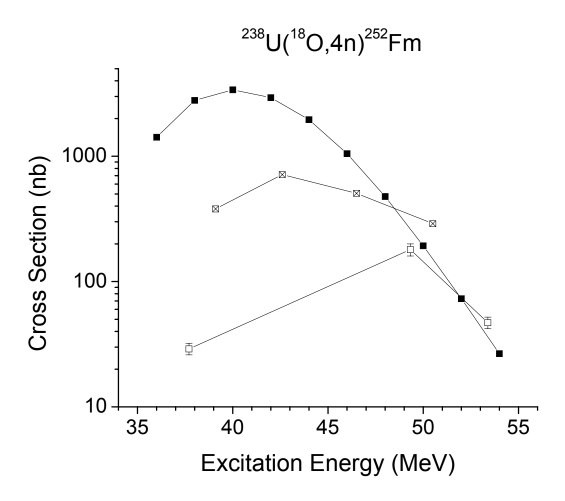


Figure 4.8: Comparison of experimental results and calculated values for the ²³⁸U(¹⁸O,4n) reaction. Solid symbols (+) represent the HIVAP code using the Reisdorf and Schädel parameters, open symbols (0) represent the experimental results of this thesis, and the open symbols with an (4) through them represent previous experimental results [Don1966].

4.3 $^{238}U + ^{22}Ne$

 256 No (E $_{\alpha}$ = 8.430(20) MeV, $t_{\frac{1}{2}}$ = 2.91 ± 0.05 s) and 257 No (E $_{\alpha}$ = 8.220(20) MeV, $t_{\frac{1}{2}}$ = 25 ± 2 s) were produced in the reaction of 22 Ne with 238 U. The Berkeley Gas-filled Separator (BGS) was used to separate the recoiling evaporation residues from the scattered beam and recoiling transfer products. The reactions specifics for the 238 U(22 Ne,xn) $^{260-x}$ No experiment are located in Table 4.9. The Coulomb barrier for the reaction was 114.9 MeV. To reduce the amount of scattering of beam that occurs with asymmetric reactions like neon on uranium, the pressure of the helium gas that enables the charge exchange collisions possible was lowered to 0.5 Torr for the entire experiment.

The ²⁶⁰No evaporation residues that were created in the hot fusion reaction in the BGS have an average momentum that was small compared with the momenta of evaporation residues resulting from cold fusion reactions. As a result, separation from transfer reaction products was difficult, and the use of a parallel plate avalanche counter (PPAC) to discriminate events that originate in the detector (alpha decay) from those that originate from beam implantations (evaporation residues) was not possible. To counter this deficiency, the beam was chopped in twenty millisecond cycles, ten milliseconds on, and ten milliseconds off. This allowed for the detection of beam events (evaporation residues) from events originating in the detector (alpha decays). Because of the high event rate in the focal plane detector during the ten millisecond beam on phase, the search for alpha events only occurred in the last five milliseconds of the beam off phase.

Table 4.9: Reaction specifics for the 238 U(22 Ne,xn) $^{260-x}$ No experiment. CYC denotes cyclotron energies, COT denotes center of target energies, and E^* denotes excitation energies.

E _{CYC} (MeV)	E _{COT} (MeV)	E* (MeV)	Current (eµA)	Dose (×10 ¹⁶)	Rutherfords	Time (s)
106.0	105.2	40.0	0.81 ± 0.07	6.1 ± 0.5	8183920	72298
110.5	109.7	44.2	0.63 ± 0.05	4.0 ± 0.3	4958640	61065
117.0	116.2	50.1	0.91 ± 0.08	4.1 ± 0.3	4505760	42829

The focal plane detector was a surface barrier silicon detector that was segmented into 32 vertical strips. The efficiency of the focal plane detector was 50 \pm 2% for alpha particles. Due to the malfunction of one of the vertical strips and the beam area being larger than the focal plane detector, only 90 \pm 5% of the evaporation residues were detected. The α -decay calibration of the focal plane detector was performed by examining the α -decay of: ¹⁹⁸Po (E $_{\alpha}$ = 6.1820(22) MeV), ¹⁹⁷Po (E $_{\alpha}$ = 6.281(4) MeV), ^{197m}Po (E $_{\alpha}$ = 6.3834(24) MeV), and ¹⁹⁶Po (E $_{\alpha}$ = 6.520(3) MeV) from the ¹¹⁴Cd(⁸⁶Kr,xn)^{200-x}Po reaction and ²⁰⁴Rn (E $_{\alpha}$ = 6.4189(25) MeV), ²⁰³Rn (E $_{\alpha}$ = 6.4992(25) MeV), ^{203m}Rn (E $_{\alpha}$ = 6.5490(25) MeV), and ²⁰²Rn (E $_{\alpha}$ = 6.6409(25) MeV) from the ¹²⁰Sn(⁸⁶Kr,xn)^{206-x}Rn reaction.

The separation efficiency was determined through the measurement of 214 Ac ($E_{\alpha} = 7.214(5)$ MeV, $t_{\%} = 8.2 \pm 0.2$ s) produced in the 197 Au(22 Ne,5n) reaction. The first step was to determine the amount of 214 Ac produced directly in the reaction. This was accomplished by placing a catcher foil directly behind a standing gold target in the BGS target chamber, collecting the recoiling products and then counting the decays of 214 Ac. A 390 μ g/cm² gold foil was used as the target and irradiated for 60 seconds by 118 MeV 22 Ne⁶⁺ at an average current of 95 enA. The foil was then quickly moved behind a silicon surface barrier detector to detect the activity of 214 Ac. The foil was located 1.36 inches from the detector, which had a diameter of 0.84 inches. It was assumed that the activity collected in the foil was in an area small enough to be considered a point source. A point source 1.36 inches from a detector of diameter 0.84 inches would have a geometric detector efficiency of $^{2+5}$ 1%. The foil was counted for a total of ten minutes in which all

of the ²¹⁴Ac decayed. The number of decays from ²¹⁴Ac were counted every 10 seconds to obtain a decay curve. The initial activity and half-life were determined from this decay curve to be $9.8 \pm 1.5 \text{ s}^{-1}$ and $8.2 \pm 0.4 \text{ s}$ (Figure 4.9). Next, the production of 214 Ac was measured in the focal plane detector. Using the same gold target and at the same energy neon projectiles at an average current of 110 enA, ²¹⁴Ac was produced and detected in the focal plane detector. Again, the number of decays from ²¹⁴Ac were counted every 10 seconds. A growth curve was drawn through the data to determine the production rate of ²¹⁴Ac as measured by the focal plane detector. The efficiency of the focal plane detector was $50 \pm 2\%$ for alpha decays. With a 390 µg/cm² gold foil, a production rate of 19 ± 3 s⁻¹ was determined (Figure 4.10). An additional experiment was performed with a thinner gold target to determine the change in efficiency with a thinner target. ²²Ne at an energy of 108 MeV and an average current of 150 enA irradiated a 100 µg/cm²gold target for approximately ten minutes. The number of ²¹⁴Ac decays was again counted every ten seconds. A curve was drawn through the data establishing the production rate for 100 $\mu g/cm^2$ target to be $19 \pm 2 s^{-1}$ (Figure 4.11). Using Equation 4.1, the BGS efficiency was calculated,

$$Eff_{BGS} = \frac{\frac{Th_{catcher}}{Th_{focal}}}{\frac{Eff_{focal}I_{focal}}{R_{catcher}}}$$

$$tword production rate from shows for the catcher fail and$$

where R is the initial activity and production rate from above for the catcher foil and focal plane measurements, Eff was the respective detector efficiencies, and I was the

beam current for each experiment. Th is the target thickness used for the particular catcher foil and focal plane detector measurements. The only experiment with the catcher foil involved a target with a thickness of 390 $\mu g/cm^2$. Focal plane measurements were made at both 390 $\mu g/cm^2$ and 100 $\mu g/cm^2$ thicknesses. To accurately calculate the efficiency of the BGS at various target thicknesses, this target thickness factor needed to be included. Using the production rates, beam currents and detector efficiencies from above and Equation 4.1, BGS efficiencies of $8 \pm 2\%$ (390 $\mu g/cm^2$) and $21 \pm 6\%$ (100 $\mu g/cm^2$) were determined for neon projectiles on gold targets. Computer simulations were run to examine the differences between BGS efficiencies for different target/projectile combinations, target thicknesses and compound nucleus velocities [Gre2002]. Extrapolation between the results of the gold target BGS efficiencies and the results of the computer simulation led to a BGS efficiency of $8 \pm 3\%$ for the 160.5 $\mu g/cm^2$ UF₄ targets and the 238 U(22 Ne,xn) reaction.

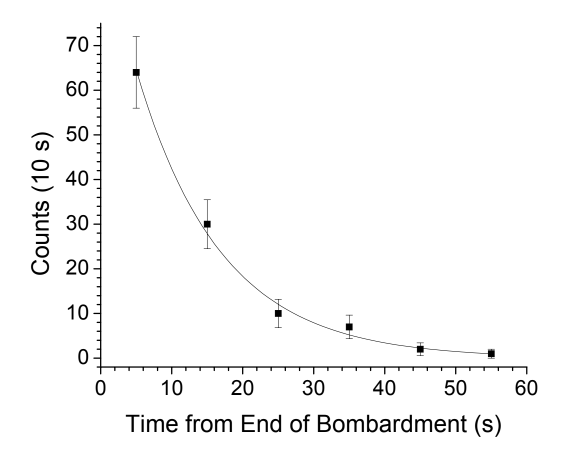


Figure 4.9: Decay data and fit from the catcher foil measurement of ^{214}Ac (t $_{\!\!\!/2}$ = $8.2\pm0.2~s$) in the reaction 118 MeV $^{22}Ne+^{197}Au$ (390 µg/cm²). An initial activity of $9.8\pm1.5~s^{-1}$ and a half-life of $8.2\pm0.4~s$ was determined.

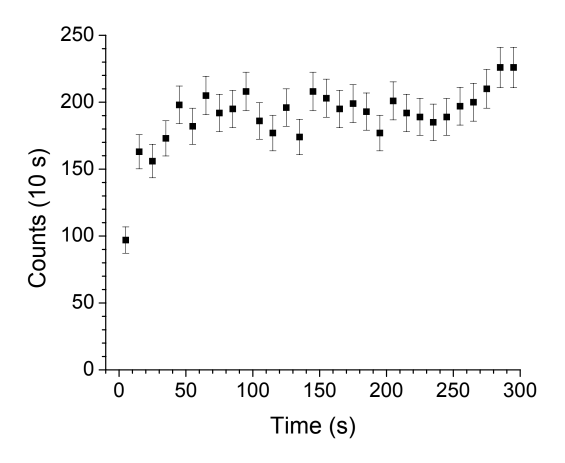


Figure 4.10: Growth data of ^{214}Ac (t $_{\!1/\!2}$ = 8.2 \pm 0.2 s) from the reaction of 118 MeV ^{22}Ne + ^{197}Au (390 $\mu g/cm^2$). A production rate of 19 \pm 3 s $^{-1}$ was determined.

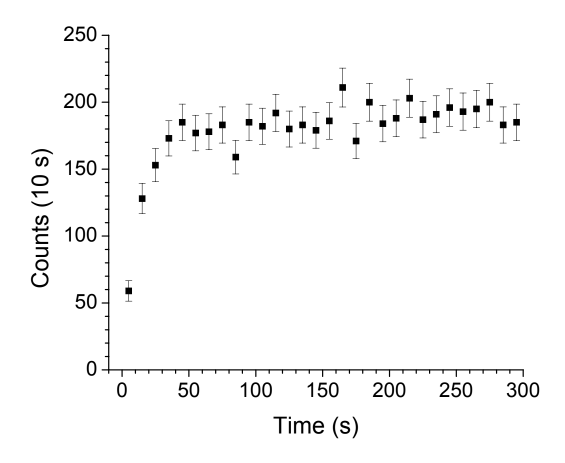


Figure 4.11: Growth data of 214 Ac ($t_{1/2}$ = 8.2 ± 0.2 s) from the reaction of 108 MeV 22 Ne + 197 Au (100 µg/cm²). A production rate of 19 ± 2 s⁻¹ was determined.

4.3.1 ²⁵⁷No

Analysis of the data obtained in the experiment was performed using the GOOSY data analysis software as described in Section 3.3. Searches were made to look for correlations between evaporation residues and 8.22 MeV, 8.27 MeV and 8.32 MeV alpha particles. After searching though the data with numerous search parameters, no correlations were found. One of the complications was the length of time between a correlated evaporation residue implantation and alpha decay on the order of the half-life of 257 No ($t_{\frac{1}{2}} = 25 \pm 2$ s). With event rates in the focal plane detector around 1000 events per second, the possibility of random correlation was high.

Assuming one correlation would have been seen, a $100 \pm 5\%$ alpha decay branching ratio, and an $8 \pm 3\%$ BGS efficiency, one-event upper limits for the production cross section were calculated for the 238 U(22 Ne,3n) reaction. Upper limit production cross sections of 4.5 nb, 6.9 nb, and 6.7 nb were calculated at excitation energies of 40.0 MeV, 44.2 MeV, and 50.1 MeV using Equations 3.10 - 3.13. A comparison of these upper limit cross sections and those obtained from the HIVAP code using the Reisdorf and Schädel parameters showed that it was reasonable not to expect any evaporation residue alpha-decay correlations. The predictions of the HIVAP code as well as the experimental upper limit production cross sections are seen in Figure 4.12.

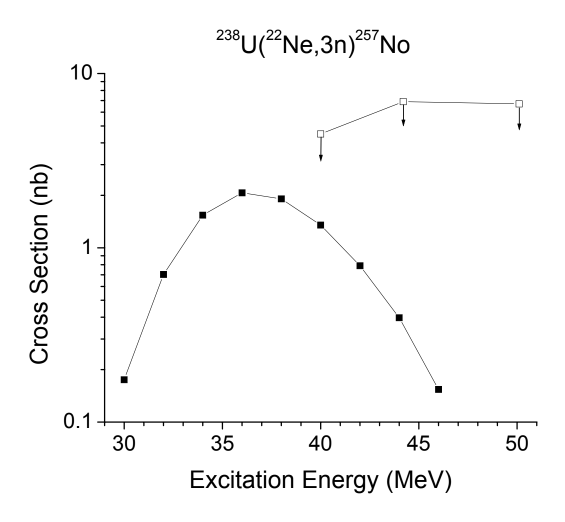


Figure 4.12: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +) and the experimental upper limit production cross sections (open symbols 0) for the 238 U(22 Ne,3n) 257 No reaction.

4.3.2 ²⁵⁶No

Unlike the search for ²⁵⁷No, the search for ²⁵⁶No was a little more successful. Using a GOOSY analysis code identical to the one in Appendix A, a search was performed to look for correlations between evaporation residue implantations and alpha decay from 256 No (E_{α} = 8.430(20) MeV, $t_{1/2}$ = 2.91 ± 0.05 s). When searching for evaporation residue alpha decay correlations, there were five main search parameters considered: decay time window, position window, evaporation residue energy window, alpha decay energy window and the beam pause/PPAC anti-coincidence window. When searching for ²⁵⁶No the following parameters were set: decay time window (0 - 30 seconds), position window (± 20 pixels, ± 1.16 mm), evaporation residue energy window (2-14 MeV), alpha decay energy (8.35-8.50 MeV), and beam pause window (15000-20000). The decay time window corresponded to the maximum amount of time allowed between an evaporation residue implantation and alpha decay. The position window corresponded to the maximum distance allowed between the correlated signals in the silicon detector. The energy windows corresponded to the energy of the evaporation residue implantations in the focal plane and the specific alpha decay energy of the isotope of interest. The beam pause window in this correlation search corresponded to the time of the twenty millisecond beam cycle in which alpha decay in the focal plane detector was considered for correlations. In this specific correlation search, only alpha decays that occurred in the last five milliseconds of the beam cycle were considered as possible correlations to evaporation residue implantations.

This correlation search was performed at all three of the experimental energies. A total of 14 correlations were seen at an excitation energy of 40.0 MeV, a total of 8

correlations were seen at an excitation energy of 44.2 MeV, and 1 event was seen at an excitation energy of 50.1 MeV. The times between the evaporation residue implantations and alpha decays were input in a program that created an input file for the MLDS program. Two component fits to the resulting decay input files were made and initial activities and total number of counts were established. The half-life for ²⁵⁶No of 2.9 seconds was fixed in the MLDS program and the second component was set to vary as a long half-life substituting for random events. In addition, all of the correlations were combined into one input file. A 2 component fit to the combined input file was performed with the first component set as a 2.9 second half-life and varied and the second component was set as a long half-life and varied. A half-life of 3.1 +2.8 seconds resulted giving weight to a successful identification of 256 No ($t_{1/2} = 2.91 \pm 0.05$ s). The results of the MLDS fits to the decay data of ²⁵⁶No at 40.0 MeV and 44.2 MeV as well as the result of the combined decay data can be seen in Table 4.10. Because only one correlation was seen at the highest excitation energy of 50.1 MeV, a one event cross section limit was calculated for the highest excitation energy. As Table 4.10 shows, a long lived component due to random correlations was present, meaning an assignment of one event at 50.1 MeV is not accurate, and therefore only a one event limit was calculated.

Using the number of counts that resulted from the MLDS fits to the correlation data, an alpha decay branching ratio of 99.5 \pm 0.1%, and a BGS efficiency of 8 \pm 3%, production cross sections were calculated for the ²³⁸U(²²Ne,4n)²⁵⁶No reaction. Production cross sections of 23 ⁺²⁰₋₁₆ nb, 14 ⁺²¹₋₉ nb, and \leq 6.8 nb were calculated at excitation energies of 40.0 MeV, 44.2 MeV, and 50.1 MeV respectively. These results

agree well with the predictions of the HIVAP code using the Reisdorf and Schädel parameters. Donets et al., performed this reaction as well and report cross sections that are similar in magnitude to the experimental cross sections observed in this thesis [Don1966]. Figure 4.13 is a combination of the experimental results of this thesis, the predictions of the HIVAP code and the experimental results of Donets et al.

Table 4.10: Results of MLDS fits to the decay curves from the correlations found in the $^{238}\text{U}(^{22}\text{Ne,4n})^{256}\text{No}$ reaction. The errors given are a result of the MLDS program. The half-life of ^{256}No ($t_{1/2}=2.91\pm0.05$ s) was fixed to eliminate the influence of random events.

Excitation Energy	Isotope	Half-life	Initial Activity	Number of Counts
40.0 MeV	²⁵⁶ No	2.9 s	$1.1^{+0.9}_{-0.7} \text{ s}^{-1}$	5 +43
44.2 MeV	²⁵⁶ No	2.9 s	$0.6^{+0.8}$ -0.2 s ⁻¹	2 +31
Combined	²⁵⁶ No	$3.1^{+2.8}_{-1.9}$ s		

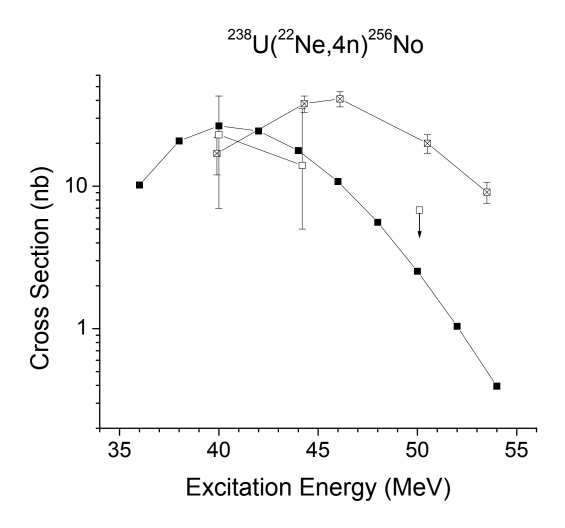


Figure 4.13: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +), the experimental production cross sections (open symbols 0), and results of Donets, et al. [Don1966] (open symbols with an 4) for the ²³⁸U(²²Ne,4n)²⁵⁶No reaction. Upper-limit cross sections denoted with an arrow.

4.4 208 Pb + 48 Ca 255 No (E $_{\alpha}$ = 8.121(6) MeV, t $_{1/2}$ = 3.1 ± 0.2 min), 254 No (E $_{\alpha}$ = 8.093(14) MeV, t $_{1/2}$ = 55 ± 3 s), 253 No (E $_{\alpha}$ = 8.010(20) MeV, t $_{1/2}$ = 1.7 ± 0.3 min), and 252 No (E $_{\alpha}$ = 8.415(6) MeV, t $_{1/2}$ = 2.30 ± 0.22 s) were all produced in the reaction of 48 Ca on 208 Pb. 48 Ca¹⁰⁺ energies between 207 MeV and 234 MeV were used in the experiment corresponding to energies at the center of the target from 204 MeV to 231 MeV and excitation energies between 12 MeV and 34 MeV. The Coulomb barrier was 217.1 MeV. The specific energies for each reaction, reaction times, average beam currents, doses, and integrated Rutherford detector counts are listed in Table 4.11.

The efficiency listed in Table 4.11 refers to the percentage of the evaporation residues that made it to the focal plane detector that were actually detected. In each of the experiments, a number of strips of the focal plane detector were not operating correctly, and therefore, an efficiency for the amount of the focal plane detector actually working during each of the reactions was determined. The focal plane detector efficiency was $50 \pm 2\%$ for alpha decay and $100 \pm 2\%$ for spontaneous fission decay.

Table 4.11: Reaction specifics for the ²⁰⁸Pb(⁴⁸Ca,xn)^{256-x}No experiment. CYC denotes cyclotron energies, COT denotes center of target energies, and E* denotes excitation energies. Eff denotes the percentage of evaporation residues that were detected by the focal plane detector. The uncertainty in the Eff value was assumed to be 5%.

E _{CYC} (MeV)	E _{COT} (MeV)	E* (MeV)	Current (eµA)	Dose (×10 ¹⁵)	Rutherford Counts	Time (s)	Eff
207.5	204.0	12.2	2.13 ± 0.11	24.9 ± 1.3	9303360	18704	80
210.0	206.5	14.2	1.61 ± 0.08	24.1 ± 1.2	8788160	23908	80
211.3	207.8	15.2	2.22 ± 0.11	19.1 ± 1.0	6890880	13812	82
212.5	208.8	16.1	0.220 ± 0.011	1.27 ± 0.06	453303	9230	64
212.5	209.0	16.3	1.66 ± 0.08	34.5 ± 1.8	12306240	33382	85
215.0	211.3	18.1	0.099 ± 0.005	0.43 ± 0.02	150966	7040	64
215.0	211.5	18.3	2.45 ± 0.12	48 ± 2	16835200	31673	87
217.5	213.8	20.2	0.161 ± 0.008	0.142 ± 0.007	48509	1414	64
218.8	215.3	21.4	0.63 ± 0.03	8.8 ± 0.4	2963936	22405	82
220.0	216.3	22.2	0.067 ± 0.003	0.43 ± 0.02	143557	10370	64
222.5	218.8	24.2	0.158 ± 0.008	0.182 ± 0.009	59036	1840	64
225.0	221.4	26.3	0.145 ± 0.007	0.46 ± 0.02	146879	5100	64
234.0	230.6	33.8	1.77 ± 0.09	19.2 ± 1.0	5632320	17384	80

BGS efficiencies for cold fusion reactions were larger than the efficiency for the 22 Ne + 238 U hot fusion reaction mentioned previously. Using Equation 3.6, the magnetic rigidity of compound nuclei and transfer products can be calculated for these two reactions. The magnetic rigidity for the hot fusion reactions was 2.00 Tm (Tesla meters) for the transfer products and 1.85 Tm for the compound nuclei. The difference between these two magnetic rigidities is about 7% and is small when compared to the difference between the rigidities of the cold fusion reaction. In the 48 Ca + 208 Pb reaction, the rigidity of the transfer products was 1.48 Tm and the rigidity of the compound nuclei was 2.12 Tm, equaling a difference of 42%. On this basis alone, a BGS efficiency approximately 5-6 times greater than the BGS efficiency used for the 22 Ne + 238 U would be expected. The actual BGS efficiency was $45 \pm 10\%$.

The experiments were run at helium pressures in the BGS of 0.74 and 0.79 Torr and a PPAC was used in one set of experiments. For the other set of experiments, the beam was pulsed 10 milliseconds on and 10 milliseconds off. Calibrations were performed by examining the products of the 165 Ho(48 Ca,xn) $^{213-x}$ Fr reaction as well as the 176 Yb(48 Ca,xn) $^{224-x}$ Th reaction. The 165 Ho(48 Ca,xn) $^{213-x}$ Fr reaction was run at 203 MeV corresponding to 50 MeV excitation energy, which led to the following products in the focal plane detector used in the calibration: 205 At (E $_{\alpha}$ = 5.902(2) MeV), 210 Rn (E $_{\alpha}$ = 6.040(2) MeV), 208 Rn (E $_{\alpha}$ = 6.1438(21) MeV), 211,210 Fr (E $_{\alpha}$ = 6.534(5) MeV, 6.543(5) MeV), and 209,208 Fr (E $_{\alpha}$ = 6.646(5) MeV, 6.641(3) MeV). The 176 Yb(48 Ca,xn) $^{224-x}$ Th reaction was also run at 203 MeV corresponding to 40 MeV excitation energy and led to the following alpha decay peaks for use in calibration: 213 Rn (E $_{\alpha}$ = 8.088(8) MeV),

 $^{221} Th~(E_{\alpha}$ = 8.472(5) MeV), and $^{215} Ra~(E_{\alpha}$ = 8.699(4) MeV). This reaction was also run at 220 MeV corresponding to 55 MeV excitation energy. The following peaks from this reaction were used for the calibration: $^{215} Ra$ and $^{217} Th~(E_{\alpha}$ = 9.250(10) MeV).

4.4.1 ²⁵⁵No and ²⁵⁴No

Disentangling the decay of ²⁵⁵No and ²⁵⁴No was difficult where the excitation functions for the 1n-exit channel and 2n-exit channel overlapped. Because of the similarities in the decay of ²⁵⁵No and ²⁵⁴No, the search for both was performed at the same time. ²⁵⁵No was seen at excitation energies from 12.1 MeV to 16.2 MeV. ²⁵⁴No was seen at excitation energies from 14.2 MeV to 26.3 MeV. The evaporation residue alpha decay correlation search parameters for ²⁵⁵No and ²⁵⁴No for the first experiment were as follows: time window (0 - 1800 seconds), position window (-20 - 15 pixels, -1.16 - 0.87 mm), evaporation residue energy window (1 – 14 MeV), alpha decay energy window (8.0 - 8.2 MeV), and PPAC window (1000 - 2500). The PPAC was used for four of the energies in which ²⁵⁵No was studied. The second experiment was performed under different conditions and a different set of correlation parameters were used: time window (0-500 seconds), position window (± 20 pixels, ± 1.16 mm), evaporation residue energy window (4 - 14 MeV), alpha decay energy window (8.0 - 8.2 MeV), and beam pause window (0-20000). It is noted that there was no effect on the number of correlations by having a shorter time window, and a smaller evaporation residue window. The event rate in the detector was small enough and the correlation time short enough to allow the use of a large beam pause window.

Figure 4.14 shows the results of the correlation search performed on the data obtained in the 215.3 MeV experiment. The top window (a.) shows all of the alpha decays that fit within the energy window and position window that were correlated to evaporation residues. The evaporation residues that correspond to the correlated alphas are located in the middle window (b.). The difference in position between the two

correlated events is found in the bottom window (c.). These windows help show that there was only one activity of interest in this particular correlation search, 254 No, the evaporation residue distribution was fairly large and covers the range between 4-10 MeV, and that the difference in position distribution was smaller than the position window used. It also appeared that some random events were present, but were accounted for when the decay curves were fit using a long component in the MLDS fit.

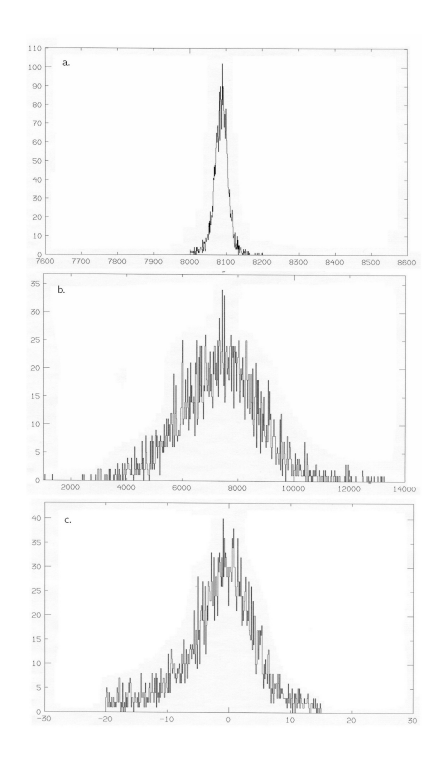


Figure 4.14: Correlated alpha decay spectrum (a.), evaporation residue spectrum (b.), and difference in position distribution (c.) for the 215.3 MeV ²⁰⁸Pb(⁴⁸Ca,2n)²⁵⁴No experiment. Energies for windows (a.) and (b.) are in keV. The difference in position distribution (c.) is in units of detector pixels.

At each excitation energy, the total number of correlations was determined from the correlation search. Then, the data were sorted according to the correlation time between the evaporation residue implantation and the alpha decay. For some of the excitation energies, only a small number of correlations ($5 \le \text{number of correlations} \le 25$) were found. In that case, the decay curve was made using a part of the MLDS program that makes input files based on only a small number of events. For the rest of the experiments (number of correlations ≤ 5), no decay curve was made. The summaries of the decay curves is found in Table 4.12. From these decay curves and those formed by the MLDS program, 2 and 3 component fits were made corresponding to the decay of ²⁵⁵No , ²⁵⁴No and a long-lived component. From the alpha decay spectra seen from the data, the presence of ²⁵⁵No and ²⁵⁴No was not guestioned and therefore their half-lives were fixed when performing the MLDS fits. The initial activities were all allowed to vary in the MLDS program. The number of counts was calculated by integrating the resulting decay curves made with the half-lives and initial activities of MLDS over the time windows used in the correlation search. Various files that corresponded to a majority of the activity of ²⁵⁵No and ²⁵⁴No were also examined with MLDS to find the corresponding experimental half-life. An experimental half-life of 169 +13 seconds was found for 255 No and an experimental half-life of 47.5 ± 0.4 seconds was found for 254 No. The summary of the MLDS fits to the data for ²⁵⁵No and ²⁵⁴No can be found in Table 4.13.

Table 4.12: Decay tables for correlations searches of evaporation residues and ²⁵⁵No and ²⁵⁴No decay. Times are in seconds and denote the start of each time bin. Counts corresponds to the number of correlations in the time bin that satisfied the correlation search parameters.

	Experiment (MeV)								
209.0			211.3	211.5	213.8	215.3	216.3	218.8	221.4
Time	Counts	Time	Counts						
0	267	0	23	1576	23	901	47	14	11
50	141	20	14	1162	21	698	32	11	10
100	85	40	7	849	10	563	25	4	5
150	43	60	4	639	5	390	19	7	5
200	27	80	7	457	10	301	16	4	2
250	10	100	3	386	10	223	11	4	4
300	19	120	0	289	2	158	9	2	2
350	13	140	1	241	2	104	5	6	2
400	6	160	3	165	0	78	2	1	1
450	7	180	2	131	1	52	3	1	0
500	8	200	3	96	1	39	1	0	0
550	5	220	1	77	0	33	6	1	0
600	3	240	3	66	0	29	1	0	0
650	3	260	1	57	1	20	3	0	0
700	1	280	0	48	1	20	3	0	0
750	3	300	0	36	0	14	1	0	0
800	1	320	1	43	2	7	1	0	0
850	3	340	1	24	1	7	1	0	1
900	2	360	0	26	0	6	1	0	0
950	5	380	0	20	0	4	0	1	0
1000	0	400	1	20	0	1	1	0	0
1050	0	420	1	13	1	1	2	0	1
1100	0	440	1	21	0	1	0	0	0
1150	0	460	0	11	1	3	2	0	0
1200	0	480	0	19	0	4	1	1	0
1250	1								
1300	0								
1350	2								
1400	0								
1450	0								
1500	0								
1550	0								
1600	1								
1650	2								
1700	3								
1750	0								

Table 4.13: MLDS results to the decay curves found in Table 4.12 (255 No $t_{\frac{1}{2}}$ = 3.1 ± 0.2 min, 254 No $t_{\frac{1}{2}}$ = 55 ± 3 s). Based on the work of [Lei1999], the half-life of 254 No was fixed at 47 s. The half-lives of 255 No and 254 No were fixed to eliminate the influence of random events.

Beam Energy	Isotope	Half-life	Initial Activity	Number of Counts
204.0 MeV	²⁵⁵ No			1.0 +2.3 -0.8
206.5 MeV	²⁵⁴ No	47 s	$0.05^{+0.05}_{-0.04} \text{ s}^{-1}$	3 +4-3
	²⁵⁵ No	186 s	$0.029^{+0.016}$ - $_{-0.012}$ s ⁻¹	8 +43
207.8 MeV	²⁵⁴ No	47 s	$0.11^{+0.09}_{-0.07} \text{ s}^{-1}$	7 +6 -5
	²⁵⁵ No	186 s	$0.09^{+0.03}_{-0.02} \text{ s}^{-1}$	25 +7 -6
208.8 MeV	²⁵⁴ No	47 s	$0.13^{+0.08}_{-0.07} \text{ s}^{-1}$	9 +5 -4
	²⁵⁵ No	186 s	$0.07^{+0.03}_{-0.02} \text{ s}^{-1}$	15 +6-5
209.0 MeV	²⁵⁴ No	47 s	$6.8 \pm 0.4 \text{ s}^{-1}$	460 ± 30
	²⁵⁵ No	186 s	$0.68^{+0.09}_{-0.08} \text{ s}^{-1}$	180 ± 20
211.3 MeV	²⁵⁴ No	47 s	$0.95^{+0.15}_{-0.14} \text{ s}^{-1}$	65 ± 10
211.5 MeV	²⁵⁴ No	47 s	$89.2 \pm 1.4 \text{ s}^{-1}$	6040 ± 90
213.8 MeV	²⁵⁴ No	47 s	$1.25^{+0.17}_{-0.16} \text{ s}^{-1}$	85 +11 -10
215.3 MeV	²⁵⁴ No	47 s	$53.7 \pm 1.1 \text{ s}^{-1}$	3640 ± 70
216.3 MeV	²⁵⁴ No	47 s	$2.5 \pm 0.2 \text{ s}^{-1}$	170 ⁺¹⁶ -15
218.8 MeV	²⁵⁴ No	47 s	$0.79^{+0.14}_{-0.12}\mathrm{s}^{-1}$	54 +9 -8
221.4 MeV	²⁵⁴ No	47 s	$0.62^{+0.12}$ -0.11 s ⁻¹	42 +8-7

Using the number of counts from the MLDS fits, half-lives based on the activity, decay branching ratios (255 No α 61.4 \pm 2.5%, 254 No α 90 \pm 4%), the data from Table 4.11 and a $45 \pm 10\%$ BGS efficiency, the cross sections were calculated using Equations 3.10 -3.13. The excitation energies, isotopes and production cross sections are listed in Table 4.14. The production cross sections were then compared to the results of the HIVAP code with the Reisdorf and Schädel parameters and to the results of previous experimental work done in Germany [Gäg1989] and Russia [Oga2000c]. The previous work on the ²⁰⁸Pb(⁴⁸Ca,xn) reaction is guite similar to the experimental results obtained in this thesis. No serious deviations exist. There does seem to be a slight cross section enhancement at lower excitation energies in the German work, but overall, the cross sections are quite similar. However, unlike the comparisons of hot fusion cross sections and predictions of the HIVAP code in which the code often closely predicted experimental results, the HIVAP code with the Reisdorf and Schädel parameters does not accurately reflect what is seen experimentally for ²⁵⁵No and is over predictive for ²⁵⁴No. This is not totally unexpected as the Reisdorf and Schädel parameters were made as a result of fits to hot fusion reactions, not cold fusion reactions. The comparison of these results can be seen in Figure 4.15 for ²⁵⁵No and Figure 4.16 for ²⁵⁴No.

Table 4.14: Production cross sections for ²⁵⁵No and ²⁵⁴No.

Excitation Energy (MeV)	Isotope	Production Cross Section (nb)
12.2	²⁵⁵ No	0.3 +0.6 -0.2
14.2	²⁵⁴ No	$0.6^{+0.8}_{-0.6}$
	²⁵⁵ No	$2.3^{\ +1.3}$ -1.0
15.2	²⁵⁴ No	1.7 +1.5 -1.3
	²⁵⁵ No	9 ± 3
16.1	²⁵⁴ No	40 ± 20
	²⁵⁵ No	100^{+50}_{-40}
16.3	²⁵⁴ No	58 ± 15
	²⁵⁵ No	33 ± 9
18.1	²⁵⁴ No	900 ± 300
18.3	²⁵⁴ No	530 ± 130
20.2	²⁵⁴ No	3500 +1000 -900
21.4	²⁵⁴ No	1900 ± 500
22.2	²⁵⁴ No	2300 ± 600
24.2	²⁵⁴ No	1700 ± 500
26.3	²⁵⁴ No	530 ± 160

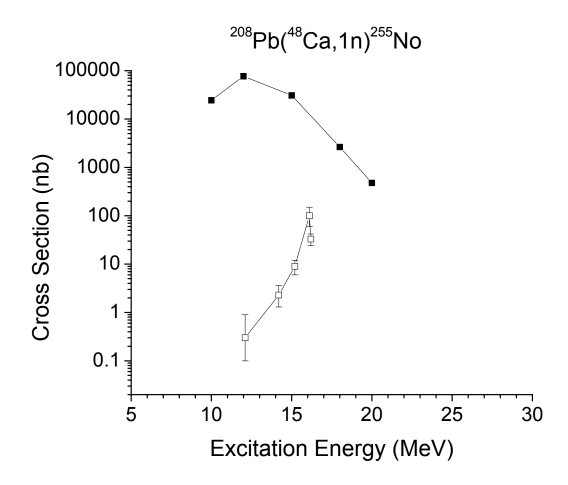


Figure 4.15: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +) and the experimental production cross sections (open symbols 0) for the ²⁰⁸Pb(⁴⁸Ca,1n)²⁵⁵No reaction.

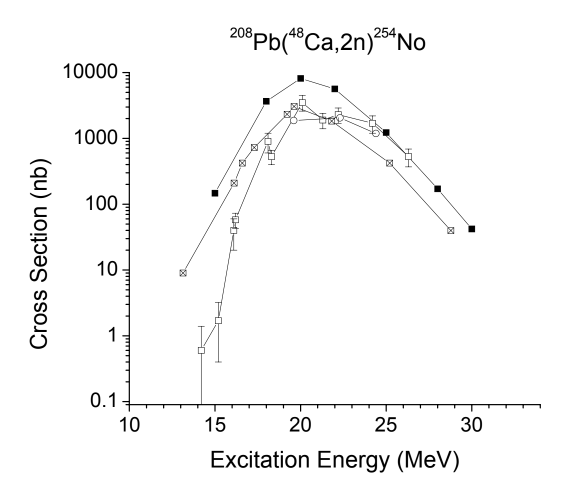


Figure 4.16: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +), the experimental production cross sections (open symbols 0), previous results of Gäggeler, et al. [Gäg1989] (open symbols with an 4), and previous results of Oganessian, et al. [Oga2000c] (open circles #) for the ²⁰⁸Pb(⁴⁸Ca,2n)²⁵⁴No reaction.

4.4.2 ²⁵³No and ²⁵²No

Correlation searches were performed to look for evaporation residue alpha decay correlations as well as evaporation residue spontaneous fission correlations. The evaporation residue alpha decay correlation search performed was similar to the correlation search performed looking for ²⁵⁵No and ²⁵⁴No with the PPAC in place. The alpha energy window was altered to look for the alpha decay of ²⁵³No (7.920 MeV – 8.201 MeV) and ^{252}No (8.37 MeV - 8.45 MeV). The evaporation residue energy window (1-14 MeV), the PPAC condition window (1000-2500), and the position difference window (-20 - 15 pixels) all remained the same. The time window between evaporation residue implantation and alpha decay was altered depending on the half-life of ²⁵³No (0 – 1000 seconds) and 252 No (0 – 25 seconds). The evaporation residue spontaneous fission correlation search differed from the search for evaporation residue alpha decay correlations in two ways. First, the energy condition for a spontaneous fission correlation search is much larger and done in channels not energy. No fission calibration was performed before the experiment and therefore no reliable energy determination was made. Second, the position window was expanded to include a much larger section of the detector. Because of the large energy signal of a fission fragment, the code is unable to calculate an accurate position determination for fission fragment signals based on the position signals present in the data. For this evaporation residue spontaneous fission correlation search, the spontaneous fission energy window was set at 500-4000 channels and the difference in position window was set at ± 250 pixels, or ± 14.5 millimeters. A total of 195 evaporation residue ²⁵³No alpha decays correlations, 6 evaporation residue ²⁵²No alpha decay correlations, and 2 evaporation residue ²⁵²No spontaneous fission

correlations were observed. A decay curve was made from the correlations observed in the ²⁵³No data with 25 second time bins. Using MLDS and the relative decay information, half-lives and initial activities for ²⁵³No and ²⁵²No were determined. The decay curve and the results of the MLDS fits can be seen in Table 4.15.

Table 4.15: Decay curves and results of MLDS fits for the correlations from the 253 No and 252 No reactions at an excitation energy of 33.8 MeV. The time bins for the decay curve are in seconds and correspond to the beginning of the time window (253 No $t_{\frac{1}{2}} = 102 \pm 18 \text{ s}$, 252 No $t_{\frac{1}{2}} = 2.30 \pm 0.22 \text{ s}$).

Time Bin	Counts	Isotope	Half-Life	Initial Activity	Number of Counts
0	35	²⁵³ No	88 ⁺⁶ -5 s	$1.55 \pm 0.13 \text{ s}^{-1}$	196 +17 -16
25	24	110	-5 5	1.33 ± 0.13 5	170 -10
50	22	²⁵² No	$3.0^{+1.2}$ -0.9 s		6
75	23	110	5.0 -0.9 5		O
100	21				
125	8				
150	13				
175	13				
200	6				
225	7				
250	6				
275	2				
300	0				
325	3 3 3				
350	3				
375					
400	1				
425	1				
450	0				
475	1				
500	1				
525	0				
550	0				
575	0				
600	1				
625	0				
650	0				
675	1				

Production cross sections were then calculated for the two reactions based on the results of the MLDS fits found in Table 4.15. The production cross section for the 208 Pb(48 Ca,3n) 253 No reaction was found to be 53 ± 14 nb at an excitation energy of 33.8 MeV and the production cross section for the ²⁰⁸Pb(⁴⁸Ca,4n)²⁵²No was found to be 1.8 ^{+1.1}_{-0.8} nb at an excitation energy of 33.8 MeV. These experimental results were then compared to the predictions of the HIVAP code using the Reisdorf and Schädel parameters. The prediction for the 3n-exit channel product, ²⁵³No, was high by a factor of four while the prediction for the 4n-exit channel product, ²⁵²No, was only high by a factor of two. These overestimations seem to get smaller as the excitation energies get higher showing that there might be a problem with the way the HIVAP parameters of Reisdorf and Schädel treat the entrance channel and its effects on the total fusion cross section. Figure 4.17 shows the comparison between the experimental data and the predictions of HIVAP for the ²⁰⁸Pb(⁴⁸Ca,3n)²⁵³No reaction and Figure 4.18 shows the comparison between the experimental data and the predictions of HIVAP for the ²⁰⁸Pb(⁴⁸Ca,4n)²⁵²No reaction.

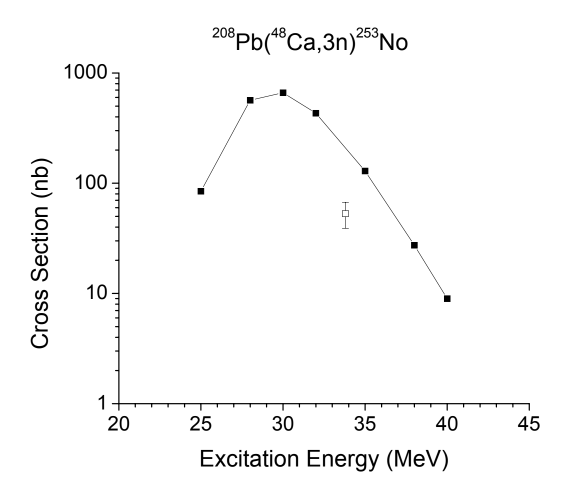


Figure 4.17: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +) and the experimental production cross sections (open symbols 0) for the ²⁰⁸Pb(⁴⁸Ca,3n)²⁵³No reaction.

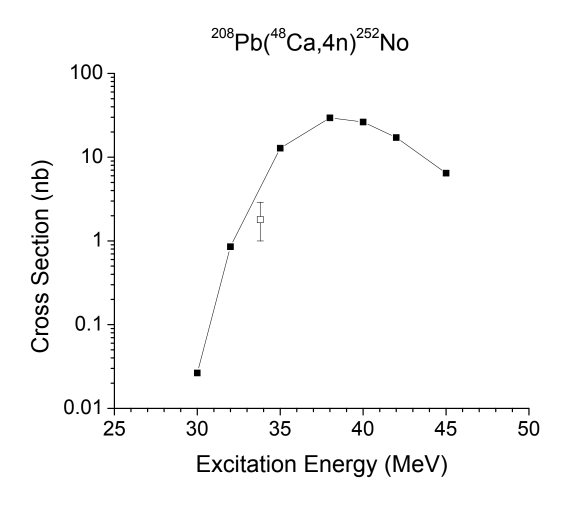


Figure 4.18: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +) and the experimental production cross sections (open symbols 0) for the ²⁰⁸Pb(⁴⁸Ca,4n)²⁵²No reaction.

$4.5 ^{208}\text{Pb} + ^{50}\text{Ti}$

 257 Rf (E $_{\alpha}$ = 8.774(8), 9.013(8) MeV, $t_{1/2}$ = 4.7 \pm 0.3s) and 256 Rf (SF, $t_{1/2}$ = 6.7 \pm 0.2 ms) were produced in the reaction of ²⁰⁸Pb and ⁵⁰Ti. The reactions specifics for the ²⁰⁸Pb(⁵⁰Ti,xn)^{258-x}Rf experiment are located in Table 4.16. The Coulomb barrier for this experiment was 239.4 MeV. The remaining experimental parameters were quite similar to previous experiments that used the BGS. The helium pressure in the BGS was held around 1.0 Torr. Approximately $90 \pm 5\%$ of the evaporation residues that made it to the focal plane detector were detected. The focal plane detector had a $50 \pm 2\%$ efficiency for detecting alpha decay and a $100 \pm 2\%$ efficiency for detecting spontaneous fission decay. The difference between the magnetic rigidities of the compound nuclei and the transfer products were similar to those obtained for the ²⁰⁸Pb(⁴⁸Ca,xn)^{256-x}No reaction, and therefore the efficiency of the BGS was $45 \pm 10\%$. A PPAC was used in the search for ²⁵⁷Rf and ²⁵⁶Rf to help discriminate between events that originated in the focal plane detector and those that occurred as a result of implantation in the focal plane detector. Calibrations for this experiment were performed by examining the decay products from the 148 Sm(50 Ti,xn) $^{198-x}$ Po reaction: 197m Po (E $_{\alpha}$ = 6.3834(24) MeV), 196 Po (E $_{\alpha}$ = 6.520(3) MeV), 195 Po (E $_{\alpha}$ = 6.609(5) MeV), 195 mPo (E $_{\alpha}$ = 6.699(5) MeV), and 194 Po (E $_{\alpha}$ = 6.843(3) MeV).

Table 4.16: Reaction specifics for the ²⁰⁸Pb(⁵⁰Ti,xn)^{258-x}Rf experiment. CYC denotes cyclotron energies, COT denotes center of target energies, and E* denotes excitation energies.

E _{CYC} (MeV)	E _{COT} (MeV)	E* (MeV)	Current (eµA)	Dose (×10 ¹⁶)	Rutherfords	Time (s)
230.0	226.0	12.9	2.78 ± 0.14	5.4 ± 0.3	19958400	37482
235.0	231.0	16.9	2.21 ± 0.11	1.27 ± 0.06	4471680	11030
237.0	233.1	18.6	1.39 ± 0.07	0.86 ± 0.04	2970880	11862
240.0	236.1	21.0	3.25 ± 0.16	2.16 ± 0.11	7297920	12776

4.5.1 ²⁵⁷Rf

Correlation searches were performed at each of the four energies studied. The first correlation search focused on looking for correlations between evaporation residues and alpha decay from ²⁵⁷Rf. The correlation search parameters were as follows: time window (0-45 seconds), evaporation residue energy window (1-14 MeV), alpha decay energy window (8.5 – 9.2 MeV), position window (± 15 pixels, ± 0.87 mm), and PPAC window (1000 - 2500). A total of 118 correlations were seen at the four different energies. No correlations were seen at the lowest energy of 226.0 MeV, 43 correlations were seen at 231.0 MeV, 29 correlations were seen at 233.1 MeV, and 46 correlations were seen 236.1 MeV. Because of the effectiveness of the BGS in separating out impurities, and because the decay region around ²⁵⁷Rf was fairly clear, all of the correlations observed were assumed to have come from evaporation residue ²⁵⁷Rf alpha decay correlations. A decay curve was made from the sum of the correlations from the four different experiments and an experimental half-life of 5.1 +0.5 =0.4 s (257 Rf $t_{1/2} = 4.7 \pm 0.3$ s) was determined using MLDS. Figure 4.19 shows the results of the correlation search for all of the experiments performed. The various decay energies of ²⁵⁷Rf are clearly seen in the top graph (a.) of Figure 4.19 and the broad distribution of the evaporation residues is seen in the middle graph (b.). Finally, the position window (c.) is clearly shown in the bottom graph and illustrates that the correlations occur within a very small separation.

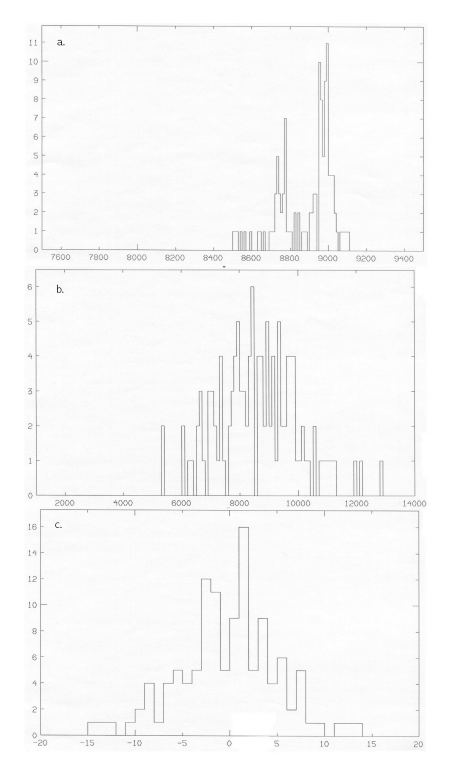


Figure 4.19: Correlated alpha decay spectrum (a.), evaporation residue spectrum (b.), and difference in position distribution (c.) for the sum of the four experiments performed in the ²⁰⁸Pb(⁵⁰Ti,1n)²⁵⁷Rf experiment. Energies for windows (a.) and (b.) are in keV. The difference in position distribution (c.) is in units of detector pixels.

A second correlation search was performed looking for alpha-alpha correlations from the decay of ²⁵⁷Rf and ²⁵³No. The correlation search parameters were as follows: time window (0 - 1000 seconds), mother alpha decay window (8.5 - 9.2 MeV), daughter alpha decay window (7.9 – 8.2 MeV), position window (± 15 pixels, ± 0.87 mm), and PPAC window (1000 - 2500). The time window and daughter alpha decay windows were chosen to look for the 1.7-minute, 8.01-MeV alpha activity of ²⁵³No. A total of 30 alpha-alpha correlations were seen between ²⁵⁷Rf and ²⁵³No. No correlations were seen at 226.0 MeV, 13 correlations at 231.0 MeV, 7 correlations at 233.1 MeV, and 10 correlations at 236.1 MeV. These numbers are close to what would be expected from a $50 \pm 2\%$ detector efficiency and an $80 \pm 5\%$ alpha decay branch for ²⁵³No. A decay curve was made from the sum of the alpha-alpha correlations and an experimental half-life of 81 $^{+13}$ ₋₁₂ s (253 No $t_{1/2}$ = 102 \pm 18 s) was determined using MLDS. Figure 4.20 shows the results of the alpha-alpha correlation search performed. The top window (a.) illustrates the decays of the daughter isotope ²⁵³No. The middle window shows the mother ²⁵⁷Rf alpha decays that were correlated to the ²⁵³No seen in the top window. The bottom window (c.) again illustrates the small distance between correlated events. The graphs clearly show the multiple decay energies present in the decay of ²⁵⁷Rf and the single decay energy of ²⁵³No.

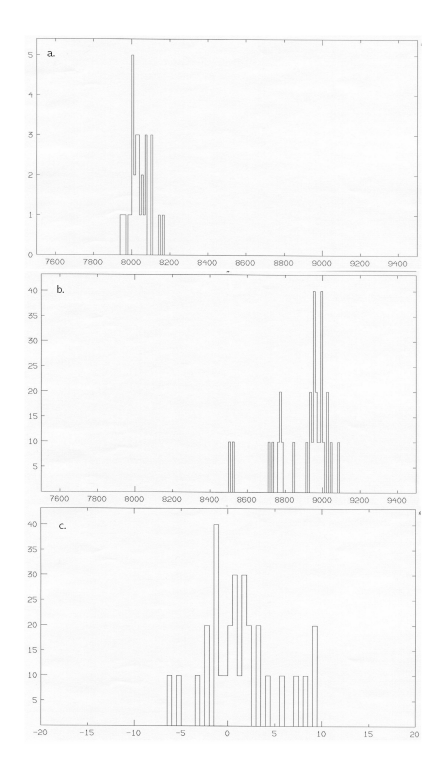


Figure 4.20: Correlated daughter alpha decay spectrum (a.), mother alpha decay spectrum (b.), and difference in position distribution (c.) for the sum of the four experiments performed in the ²⁰⁸Pb(⁵⁰Ti,1n)²⁵⁷Rf experiment. Energies for windows (a.) and (b.) are in keV. The difference in position distribution (c.) is in units of detector pixels.

Using the number of correlations found at each of the individual energies, a $79.6 \pm 2.0\%$ alpha decay branching ratio, and a $45 \pm 10\%$ BGS efficiency, the production cross sections were calculated for the ²⁰⁸Pb(⁵⁰Ti,1n)²⁵⁷Rf reaction at the four energies studied. The production cross sections obtained at the associated excitation energies were: ≤ 0.086 nb (12.9 MeV), 16 ± 4 nb (16.9 MeV), 16 ± 5 nb (18.6 MeV), and 10 ± 3 nb (21.0 MeV). The cross section at 12.9 MeV is a one event upper limit cross section as no correlations that matched the search parameters were found. These cross sections were compared with predictions from the HIVAP code using the Reisdorf and Schädel parameters. As was seen with the ²⁰⁸Pb(⁴⁸Ca,1n)²⁵⁵No reaction, the 1n-exit channel predictions of HIVAP greatly overestimate the production cross sections. The experimental cross sections were smaller than the predictions by a couple of orders of magnitude at the lower energies. It also appears that the HIVAP code incorrectly predicts the center of the excitation function, usually to lower excitation energies than what is seen experimentally. The experimental cross sections were also compared to previous experimental ²⁰⁸Pb(⁵⁰Ti,1n)²⁵⁷Rf results of Heβberger, et al. [Heβ1997]. These experimental cross sections are a lot closer in magnitude than the predictions of HIVAP. There still appears to be a difference in the center of the excitation function between the results of Heßberger and this thesis. It appears the maximum of the 1n-exit channel excitation function appears around 16 MeV for Heßberger whereas the maximum appears around 18-19 MeV for the results of this thesis. Figure 4.21 shows the comparison of the predictions from HIVAP and the experimental results for the ²⁰⁸Pb(⁵⁰Ti,1n)²⁵⁷Rf reaction.

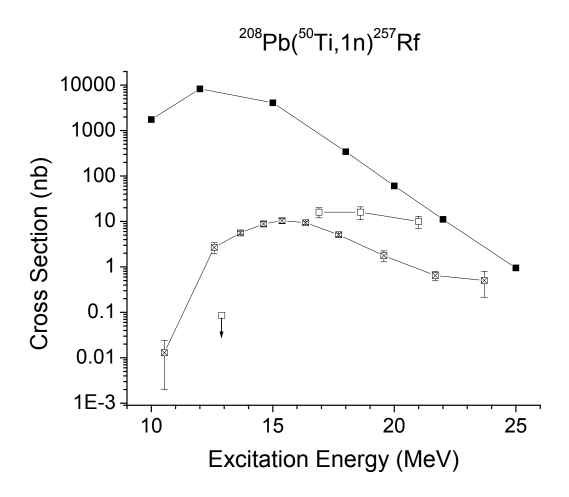


Figure 4.21: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +), the experimental production cross sections (open symbols 0), and the previous results of Heßberger, et al. [He β 1997] (open symbols with an 4) for the $^{208}\text{Pb}(^{50}\text{Ti},1n)^{257}\text{Rf}$ reaction.

4.5.2 ²⁵⁶Rf

Correlations searches were performed looking for the spontaneous fission decay of ²⁵⁶Rf in the attempt to calculate cross sections for the 2n-exit channel from the ²⁰⁸Pb(⁵⁰Ti,2n) reaction. Spontaneous fission correlation searches differ from evaporation residue alpha decay correlation searches in two ways. The first difference is that the position window is larger when searching for fission correlations. Fission products are more energetic than alpha decays and therefore are more likely to travel farther in the silicon detector. The large amount of energy deposited by the fission products also makes position determination more difficult. The focal plane detector was setup so that high energy fission products occurring near the top or bottom of a strip were likely to saturate the signal making accurate position determinations difficult. Therefore a larger position window is used when looking for evaporation residue spontaneous fission correlations. The second difference lies in the energy calibrations for the higher energy fissions. Spontaneous fission sources are not used in the calibration of the focal plane detector to eliminate the possibility of background fission activity in the detector. Without an appropriate energy calibration, the spontaneous fission energy window is in channel numbers and not keV. Aside from these two differences, the rest of the correlation search procedure was similar to those already performed.

When looking for the 256 Rf produced in the 208 Pb(50 Ti,2n) reaction, the following correlation search parameters were set. Time window (0 – 70 milliseconds), evaporation residue energy window (1 – 14 MeV), spontaneous fission energy window (1650 – 4000 channels), position window (± 250 pixels, ± 14.5 mm), and PPAC condition window (± 250). These correlation search parameters were used to find 4 evaporation

residue fission correlations at an excitation energy of 16.9 MeV, 5 evaporation residue fission correlations at an excitation energy of 18.6 MeV, and 77 evaporation residue fission correlations at an excitation energy of 21.0 MeV. The trend to higher numbers of correlations was consistent with the increase in excitation energy moving from an area where the 1n-exit channel was more prominent to an area where the 2n-exit channel was more prominent. A sum of all of the correlations between evaporation residues and spontaneous fissions is shown in Figure 4.22.

The specific times between evaporation residue implantation and spontaneous fission were taken for each energy and made into decay curves for input into the decay curve fitting program MLDS. Initial activities as well as half-lives and the total number of counts at each excitation energy were obtained from the MLDS program. An experimental half-life of $7.0^{+0.7}$ -0.6 ms (256 Rf $t_{1/2} = 6.7 \pm 0.2$ ms) was obtained for 256 Rf.

Using the number of counts obtained from MLDS, a detection efficiency of $100 \pm 2\%$, a BGS efficiency of $45 \pm 10\%$, and a 98^{+2} - $_7\%$ spontaneous fission branching ratio, cross sections for the $^{208}\text{Pb}(^{50}\text{Ti},2\text{n})^{256}\text{Rf}$ reaction were obtained as follows: $0.6^{+0.5}$ - $_{0.3}$ nb (16.9 MeV), $1.1^{+0.8}$ - $_{0.5}$ nb (18.6 MeV), and 6.7 ± 1.8 nb (21.0 MeV). These cross sections are compared to the results of Heßberger, et al. [Heß1997], and compare fairly well. With the energy spread used, it was again easy to see the energy differences on the low energy side of the excitation functions. This discrepancy can only be explained by a differing energy between the two machines used to create the ^{50}Ti beams used in these experiments. The 2n-exit channel results were also compared to the predictions of the HIVAP code using the input parameters of Reisdorf and Schädel and

appear to be just as different from the code as the rest of the cold fusion reactions already shown. The cross section comparison can be seen in Figure 4.23.

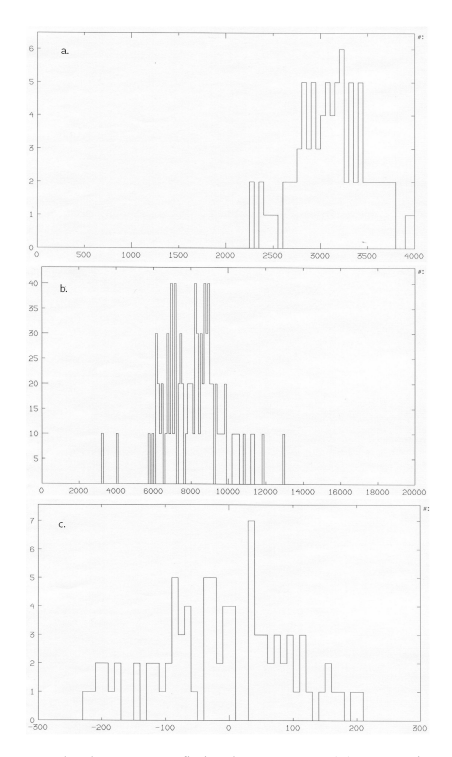


Figure 4.22: Correlated spontaneous fission decay spectrum (a.), evaporation residue spectrum (b.), and difference in position distribution (c.) for the sum of the three experiments performed in the ²⁰⁸Pb(⁵⁰Ti,2n)²⁵⁶Rf experiment. The top graph (a.) is in channels whereas the energy for window (b.) is in keV. The difference in position distribution (c.) is in units of detector pixels.

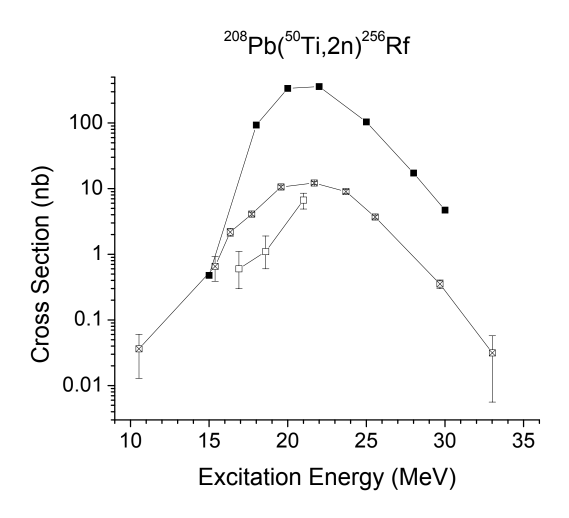


Figure 4.23: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +), the experimental production cross sections (open symbols 0), and the previous results of Heßberger, et al. [He β 1997] (open symbols with an 4) for the 208 Pb(50 Ti,2n) 256 Rf reaction.

$4.6 \quad ^{208}\text{Pb} + ^{51}\text{V} \rightarrow ^{257}\text{Db} + 2\text{n}$

The ²⁰⁸Pb(⁵¹V,2n)²⁵⁷Db reaction was examined in the attempt to study a possible production route of dubnium for chemical study. This reaction was chosen and would be compared with a similar reaction, ²⁰⁹Bi(⁵⁰Ti,2n)²⁵⁷Db discussed subsequently, which produced the same compound nucleus.

⁵¹V¹¹⁺ at energy of 255.0 MeV from the cyclotron lost approximately 5 MeV in the carbon windows and half of the target leaving 250.0 MeV in the center of the target which corresponded to an excitation energy of 24.9 MeV. The Coulomb barrier was 250.7 MeV. The experiment lasted 38535 seconds delivering a total beam dose of $(2.90 \pm 0.14) \times 10^{16}$ particles for an average current of 1.33 ± 0.07 eµA. A total of 10375000 Rutherford events were collected during the experiment. The compound nucleus recoils travel through 1 Torr of helium in the gas-filled chamber and had a magnetic rigidity of 2.05 Tm whereas the transfer products of the reaction had a magnetic rigidity of 1.48 Tm giving a BGS efficiency of $45 \pm 10\%$. No PPAC was used in this particular experiment, but because of the short half-life of 257 Db (E_a = 8.970(20) MeV, $t_{1/2} = 1.3^{+0.5}$ -0.3 s), it wasn't a concern. The experiment was calibrated using the ¹⁵⁴Sm(⁵¹V,xn)^{205-x}At reaction at an excitation energy of 75.8 MeV which would lead to the emission of between five and eight neutrons. The following activities were used in the calibration: 200 Po (E_{α} = 5.8619(18) MeV), 199m Po (E_{α} = 6.059(3) MeV), 198 Po (E_{α} = 6.1820(22) MeV, $^{200\text{m}}$ At (E_{\alpha} = 6.5374(23) MeV), 199 At (E_{\alpha} = 6.643(3) MeV), 198 At (E_{\alpha} = 6.754(4) MeV), and 198m At (E_{α} = 6.856(4) MeV).

Once the data was collected, correlation searches were performed looking for both evaporation residue alpha decay correlations as well as alpha-alpha decay correlations. The alpha decay daughter of 257 Db is 253 Lr ($E_{\alpha} = 8.800(20)$ MeV, $t_{\frac{1}{2}} = 1.3^{+0.6}$ -0.3 s). With these decay half-lives and decay energies as a guide the following correlation parameters were set looking for the evaporation residue alpha decay correlations: time window (0 – 15 seconds), position window (± 15 pixels, ± 0.87 mm), evaporation residue energy window (2 - 14 MeV), alpha decay energy window (8.90 - 9.25 MeV). The following correlation parameters were set for the search for alpha-alpha correlations: time window (0 - 15 seconds), position window (± 15 pixels, ± 0.87 mm), mother alpha energy window (8.90 - 9.25 MeV), daughter alpha energy window (8.65 - 8.87 MeV).

A total of 10 evaporation residue alpha decay correlations were seen as well as 5 alpha-alpha correlations. The number of alpha-alpha correlations is expected considering the branching ratio for alpha decay for 253 Lr is $98 \pm 2\%$ and the detection efficiency for alpha decay in the focal plane detector was $50 \pm 2\%$. From these efficiencies, five alpha-alpha decay correlations would be expected and were seen. Decay curves were made from the correlation data and analyzed using MLDS. Half-lives and initial activities were determined from the MLDS fits and used to calculated the cross sections. Experimental half-lives of 1.1 ± 0.2 s (257 Db t_{1/2} = $1.3^{+0.5}$ -0.3 s) for 257 Db and $0.9^{+0.4}$ -0.3 s (254 Lr t_{1/2} = $1.3^{+0.6}$ -0.3 s) for 253 Lr were determined.

Using the number of evaporation residue alpha decay correlations, a BGS efficiency of $45 \pm 10\%$, a $95 \pm 5\%$ evaporation residue detection efficiency, and a $82 \pm 11\%$ alpha-decay branching ratio for 257 Db, a production cross section was

calculated for the one experimental energy. A production cross section of 1.4 ^{+0.7}-_{0.6} nb was calculated for an excitation energy of 24.9 MeV. As the experiment only considered one experimental energy, an assessment of the behavior of the excitation function was difficult. Nonetheless, this experimental point was compared with the predictions of the HIVAP code using the Reisdorf and Schädel parameters. Again, the HIVAP predictions are not similar to the cross sections seen experimentally. The comparison of the predictions with the experimental cross section can be seen in Figure 4.24.

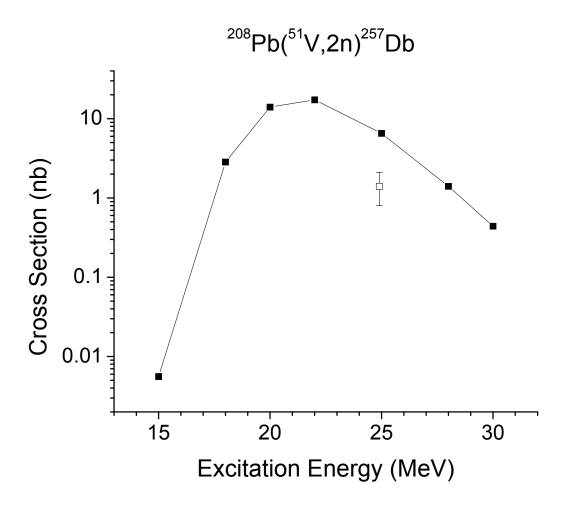


Figure 4.24: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +), the experimental production cross sections (open symbols 0) for the 208 Pb(51 V,2n) 257 Db reaction.

$4.7 \quad ^{209}\text{Bi} + ^{50}\text{Ti}$

The ²⁰⁹Bi + ⁵⁰Ti reaction was examined as another possible production route for possible chemistry experiments with dubnium. The experiment was also examined to look at the effects of different target and projectile combinations forming the same compound nucleus. Two separate experiments were performed for this reaction at two different energies. The Coulomb barrier for this reaction was 253.2 MeV.

The first experiment was performed with ⁵⁰Ti¹²⁺ beams at an energy of 235 MeV, which then lost approximately 4 MeV as it passed through the carbon windows and half of the bismuth target leaving 231.2 MeV in the center of the target. This corresponded to a compound nucleus excitation energy of 15.0 MeV.

The experiment lasted 13413 seconds at an average beam current of 2.87 ± 0.15 eµA which corresponded to a total beam dose of $(2.00 \pm 0.10) \times 10^{16}$. A total of 6091472 Rutherford scattered beam events were recorded during the experiment. The magnetic rigidity of the compound nucleus recoils were 2.10 Tm whereas the magnetic rigidity of the transfer products was 1.48 Tm. This led to a BGS efficiency of $45 \pm 10\%$. No PPAC was used in this first experiment. About $90 \pm 5\%$ of the evaporation residues making it to the focal plane detector were actually detected. This first experiment had no calibration reaction.

The second experiment was run at a slightly higher energy. 237 MeV ⁵⁰Ti¹²⁺ lost approximately 3.5 MeV leaving 233.5 MeV at the center of the target which corresponded to a compound nucleus excitation energy of 16.9 MeV.

The second experiment lasted a total of 23510 seconds at an average beam current of 1.68 ± 0.09 eµA giving a total beam dose of $(2.05 \pm 0.11) \times 10^{16}$. A total of 6293120

Rutherford scattered beam events were recorded. The magnetic rigidities of the evaporation residues and transfer products in the BGS remained 2.10 Tm and 1.48 Tm respectively, leaving the BGS efficiency at $45 \pm 10\%$. A PPAC detector was used in this second experiment to discriminate between events originating in the focal plane detector and those recorded as implantation events. Again, approximately $90 \pm 5\%$ of the evaporation residue events making it to the focal plane detector were actually detected.

The focal plane detector for the second experiment was calibrated using the nat Dy(50 Ti,xn) $^{214-x}$ Ra reaction. The most prominent isotopes of natural dysprosium are 162 Dy, 163 Dy, and 164 Dy. At an excitation energy of approximately 55-60 MeV, the 4n-and 5n-exit channels were the preferred neutron evaporation channels. The focal plane detectors were calibrated using the following isotopes: 205 Rn (E $_{\alpha}$ = 6.262(3) MeV), 204 Rn (E $_{\alpha}$ = 6.4189(25) MeV), 208,209 Fr (E $_{\alpha}$ = 6.641(3) MeV, 6.646(5) MeV), 209,210 Ra (E $_{\alpha}$ = 7.008(5) MeV, 7.019(5) MeV), and 208 Ra (E $_{\alpha}$ = 7.133(5) MeV).

4.7.1 ²⁵⁸Db and ²⁵⁷Db

The 258 Db (E $_{\alpha}$ = 9.172(15) MeV, $t_{1/2}$ = 4.4 $^{+0.9}$ -0.6 s) produced in this reaction was produced at excitation energies corresponding to the supposed maximum in the 209 Bi(50 Ti,1n) excitation function. Correlation searches were aimed at finding the evaporation residue alpha decay correlations of 258 Db as well as possible evaporation residue electron capture spontaneous fission correlations of 258 Rf (SF, $t_{1/2}$ = 12 ± 2 ms)

The first correlation search focused on looking for the evaporation residue alpha decay correlations for 258 Db at 231.2 MeV. In this particular experiment, no PPAC was used and no energy calibration was present making the search for decays somewhat difficult. Nonetheless, a broad energy window was used ($\sim 8-10$ MeV), with the common position window and no positive correlations were seen. A possible correlation was seen, but without a proper energy calibration, a positive identification was impossible. A one event upper limit production cross section of 0.33 nb was calculated for 258 Db at an excitation energy of 15.0 MeV.

The second correlation search again looked for the evaporation residue alpha decay correlations of 258 Db, this time at 233.5 MeV. Identification of positive correlations was made easier by the use of the PPAC and a proper calibration. The following correlation search parameters were used: time window (0 – 45 seconds), position window (± 15 pixels, ± 0.87 mm), evaporation residue energy window (1-14 MeV), alpha decay energy window (8.8 - 9.5 MeV), and PPAC window (1000 - 2500). A total of 10 correlations was seen at 233.5 MeV. The experimental half-life of the activity was determined from an MLDS fit to the lifetimes of the 10 correlations to be $2.9^{+1.0}$ - $_{0.8}$ seconds. The possibility that some of the events could have resulted from

 257 Db (E_{α} = 8.970(20) MeV, $t_{\frac{1}{2}}$ = 1.3 $^{+0.5}$ - $_{0.3}$ s) was small. On one side, the excitation energy is more favorable to the 1n-exit channel versus the 2n-exit channel, and two, no correlations were seen at the most prominent energy for 257 Db at 8.970 MeV reducing the probability that decays at the other energies were possible. With a 50 \pm 2% alpha decay detection efficiency and a 67 $^{+5}$ - $_{9}$ % alpha decay branching ratio, a production cross section of 3.1 $^{+1.5}$ - $_{1.3}$ nb was calculated at an excitation energy of 16.9 MeV. Using an 82 \pm 11% alpha decay branching ratio, a one event upper limit production cross section of 0.25 nb was calculated for the 2n-exit channel 257 Db.

The third correlation search focused on looking for spontaneous fissions in correlation with evaporation residues. The spontaneous fissions would arise from the decay of 258 Rf, the electron capture decay daughter of 258 Db. The following search parameters were used: time window (0 – 45 seconds), position window (± 250 pixels, ± 14.5 mm), evaporation residue energy window (1 – 14 MeV), spontaneous fission channel window (1650 - 4000), and PPAC window (1000 - 2500). A total of 8 evaporation residue spontaneous fission correlations were seen using these parameters. Taking into account a $50 \pm 2\%$ alpha detection efficiency, a 33^{+9} - $_5\%$ electron capture branching ratio versus a 67^{+5} - $_9\%$ alpha decay branching ratio, a $100 \pm 2\%$ spontaneous fission detection efficiency, and an $87 \pm 11\%$ spontaneous fission branching ratio in 258 Rf, we should have expected 8.7 spontaneous fissions. This results gives further support to claim that only the 1n-exit channel product, 258 Db was seen, and not 257 Db the 259 Deceived the spontaneous first only the 1n-exit channel product.

These production cross sections and upper limit cross sections were compared to previous experiments of Heßberger et al. [Heß2001a]. The comparisons with the available experimental data are somewhat difficult because of the fact that an incomplete excitation function was measured. Therefore, having only a few experimental points makes comparison difficult. However, it does appear that an energy shift in the 1n-exit channel excitation function is indeed present as it has been in the experiments already discussed. The absence of correlations at 15.0 MeV excitation energy is questionable though and could be the result of faulty experimental equipment. Without additional data points however, a more accurate comparison can not be performed. The experimental points were also compared to the results of predictions by the HIVAP code using the Reisdorf and Schädel parameters. The HIVAP code predictions again predict cross sections that are higher than those seen experimentally at these low excitation energies. Figure 4.25 and Figure 4.26 shows the results of the ²⁰⁹Bi(⁵⁰Ti,1n)²⁵⁸Db and ²⁰⁹Bi(⁵⁰Ti,2n)²⁵⁷Db reactions along with other experimental results and the predictions of HIVAP.

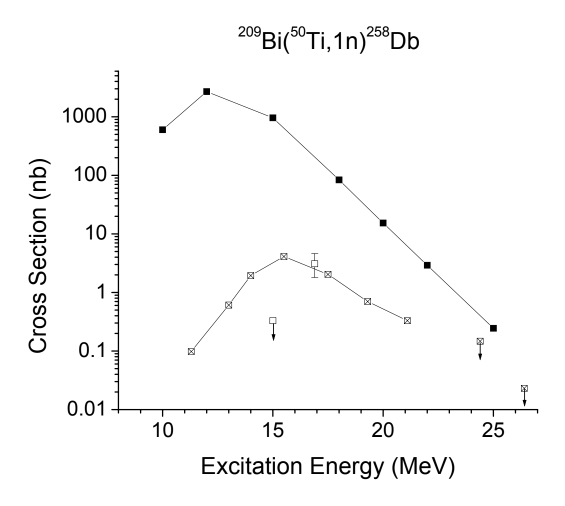


Figure 4.25: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +), the experimental production cross sections (open symbols 0), and the previous results of Heßberger, et al. [Heβ2001a] (open symbols with an 4) for the 209 Bi(50 Ti,1n) 258 Db reaction.

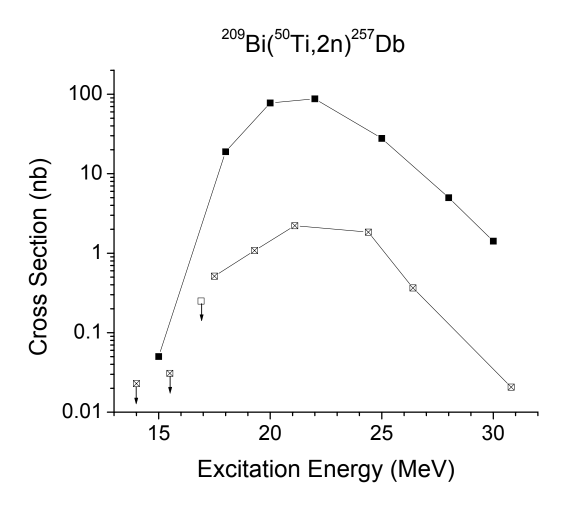


Figure 4.26: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +), the experimental production cross sections (open symbols 0), and the previous results of Heßberger, et al. [Heβ2001a] (open symbols with an 4) for the 209 Bi(50 Ti,2n) 257 Db reaction.

4.8 $^{209}\text{Bi} + ^{51}\text{V} \rightarrow ^{258}\text{Sg} + 2\text{n}$

 258 Sg (SF, $t_{1/2} \approx 2.9$ ms) was produced in the reaction of 209 Bi and 51 V in the attempt to study the effect an odd number of protons in the target and projectile would have on production cross sections. Measurements had already been made on the even proton number reaction 208 Pb + 50 Ti. Adding a proton to both the target and projectile gave the vanadium on bismuth reaction. Studying the effects the odd protons have on the production cross sections would help determine future production routes to the heavy elements.

The reactions specifics for the 209 Bi(51 V,2n) 258 Sg experiment are located in Table 4.17. The Coulomb barrier for this reaction was 253.2 MeV. The compound nuclei in these reactions had a magnetic rigidity of 2.06 Tm whereas the transfer products had a magnetic rigidity of 1.48 Tm. These rigidities corresponded to a BGS efficiency of $45 \pm 10\%$. A PPAC was used in this experiment as in others to discriminate between evaporation residue implantations in the focal plane detector and fission events from 258 Sg. The event rate in the focal plane detector was small for this experiment, so it was assumed that $100 \pm 2\%$ of the evaporation residues that made it to the focal plane detector were detected. No spontaneous fission calibration had been performed prior to the experiment as long-lived spontaneous fission isotopes have a tendency to build up on the detector when performing calibration experiments.

Table 4.17: Reaction specifics for the ²⁰⁹Bi(⁵¹V,2n)²⁵⁸Sg experiment. CYC denotes cyclotron energies, COT denotes center of target energies, and E* denotes excitation energies.

E _{CYC} (MeV)	E _{COT} (MeV)	E* (MeV)	Current (eµA)	Dose (×10 ¹⁷)	Rutherfords	Time (s)
249.0	245.4	20.3	2.80 ± 0.14	1.37 ± 0.07	41446400	93943
252.0	248.4	22.7	2.68 ± 0.14	2.13 ± 0.11	62864000	152673
255.0	251.4	25.2	1.99 ± 0.10	2.26 ± 0.12	65129600	217916

Once all of the data had been collected, a detailed correlation search began looking for evaporation residue spontaneous fission correlations. Because the half-life of 258 Sg is approximately 2.9 milliseconds and decays primarily (~100%) through spontaneous fission [He β 1997], the number of expected random correlations was extremely small (< 0.007 for each evaporation residue event). The following conditions were used to perform the correlation search: time window (0 – 50 milliseconds), position window (\pm 250 pixels, \pm 14.5 mm), evaporation residue channel window (0 – 4000), spontaneous fission channel window (1650 – 4000), and PPAC window (1000 – 2500). Channel windows were used because no spontaneous fission energy calibration had been performed before the experiment. The window sizes were chosen based on previous experimental evidence for the appearance of fission events in the data.

A total of nine evaporation residue spontaneous fission correlation events were found at the three different energies. In the data, all of the correlations were seen with an evaporation residue event followed by a spontaneous fission as the next event. Table 4.18 contains a summary of the results of the correlation search for ²⁵⁸Sg.

Table 4.18: Summary of evaporation residue spontaneous fission correlations from the ²⁰⁹Bi(⁵¹V,2n)²⁵⁸Sg reaction. The energy "E" is listed as the center of target beam energy. "Strip" corresponds to the detector strip in which the event occurred. "dt" and "dp" correspond to the differences in time and position between the evaporation residue "EVR" and spontaneous fission "SF" events which are listed in channels.

E (MeV)	Date	Time	Strip	EVR (chan)	SF (chan)	dt (ms)	dp (mm)
245.4	10/15/01	08:35	14	3022	3111	6.005	7.1
248.4	10/13/01	11:12	3	2556	2940	4.019	3.4
	10/13/01	12:14	2	2363	3234	2.873	9.3
	10/13/01	21:56	12	2203	3113	7.053	6.7
	10/13/01	23:01	11	2479	2667	1.632	2.8
	10/14/01	12:27	17	2190	3585	2.582	9.9
251.4	10/10/01	07:03	8	2342	2541	4.779	1.2
	10/10/01	22:51	5	2422	3019	3.709	1.6
	10/11/01	05:40	4	2019	2980	2.924	9.8

The channels for the evaporation residues and spontaneous fissions listed in Table 4.18 are consistent with evaporation residue spontaneous fission correlations done previously (Section 4.5.2, Figure 4.22). The correlated evaporation residue channels corresponded to energies between 8 and 12 MeV, well within the range of evaporation residues seen in previous experiments.

The lifetimes of the correlation events was taken and fit using MLDS to establish an experimental half-life for 258 Sg ($t_{1/2} \approx 2.9$ ms). One previous experiment has listed a half-life for 258 Sg of $2.9^{+1.3}$ - $_{0.7}$ milliseconds [He β 1997]. An experimental half-life of $2.7^{+0.9}$ - $_{0.7}$ milliseconds was obtained from the MLDS fit to the data.

Production cross sections were calculated from the number of correlations seen, a 100% fission detection efficiency and a 100 \pm 5% spontaneous fission branching ratio to be 14 $^{+32}$ - $_{12}$ pb, 50 $^{+30}$ - $_{20}$ pb, and 26 $^{+26}$ - $_{15}$ pb corresponding to compound nucleus excitation energies of 20.3 MeV, 22.7 MeV, and 25.2 MeV. These experimental values were compared to experimental data from GSI [He β 2001b] and the results of the HIVAP code using the Reisdorf and Schädel parameters. The production cross section comparison for the 209 Bi(51 V,2n) 258 Sg reaction can be seen in Figure 4.27. The small number of correlations leads to large error bars and a difficulty in comparing the experimental results presented in Figure 4.27. The experimental results are however consistent in the fact that they are both smaller by more than an order of magnitude than the predictions of the HIVAP code.

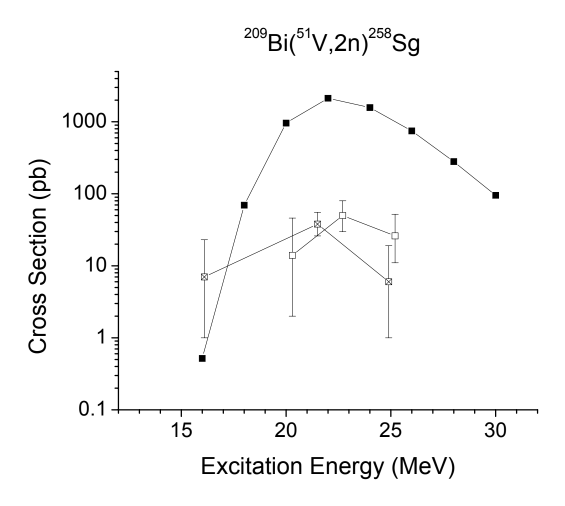


Figure 4.27: Predictions of the HIVAP code using the Reisdorf and Schädel parameters (solid symbols +), the experimental production cross sections (open symbols 0), and the previous results of Heßberger, et al. [Heβ2001b] (open symbols with an 4) for the $^{209}\mathrm{Bi}(^{51}\mathrm{V},2n)^{258}\mathrm{Sg}$ reaction.

Cross sections from this reaction of a projectile and target with an odd proton was smaller than the similar even proton projectile and target reaction ²⁰⁸Pb(⁵⁰Ti,2n)²⁵⁶Rf by almost a factor of 200. Obviously the odd projectile and targets have an effect on the production cross section reducing it by a factor of about 200, but how that effect manifests itself is still unknown.

5 Discussion

5.1 Cross sections

Tables 5.1 and 5.2 contain a summary of the experimental production cross sections of this thesis. Each reaction is listed with the beam energy at the center of the target, the excitation energy at the center of the target, and the production cross section. The hot fusion reaction summary is seen in Table 5.1 and the cold fusion reaction summary is found in Table 5.2.

Table 5.1: Summary of experimental hot fusion production cross sections. Cross sections without errors are upper limit cross sections only.

Reaction	Energy (MeV)	E* Energy (MeV)	Cross Section
248 Cm(15 N,3n) 260 Lr	75.9	35.2	$2.3 \pm 1.2 \text{ nb}$
248 Cm(15 N,3n) 260 Lr	78.8	37.9	$2.8 \pm 1.4 \text{ nb}$
248 Cm(15 N,4n) 259 Lr	75.9	35.2	$27^{+17}_{-14}\mathrm{nb}$
248 Cm(15 N,4n) 259 Lr	78.8	37.9	$90 \pm 40 \text{ nb}$
$^{238}\text{U}(^{18}\text{O},4\text{n})^{252}\text{Fm}$	82.4	37.7	29 ± 3 nb
$^{238}\text{U}(^{18}\text{O},4\text{n})^{252}\text{Fm}$	94.9	49.3	$180 \pm 20 \text{ nb}$
$^{238}\text{U}(^{18}\text{O},4\text{n})^{252}\text{Fm}$	99.3	53.4	$47 \pm 5 \text{ nb}$
²³⁸ U(²² Ne,3n) ²⁵⁷ No	105.2	40.0	≤ 4.5 nb
$^{238}\text{U}(^{22}\text{Ne,3n})^{257}\text{No}$	109.7	44.2	\leq 6.9 nb
$^{238}\text{U}(^{22}\text{Ne,3n})^{257}\text{No}$	116.2	50.1	≤ 6.7 nb
$^{238}U(^{22}Ne,4n)^{256}No$	105.2	40.0	$23^{+20}_{-16} \text{ nb}$
$^{238}\text{U}(^{22}\text{Ne,4n})^{256}\text{No}$	109.7	44.2	14 ⁺²¹ -9 nb
$^{238}\text{U}(^{22}\text{Ne,4n})^{256}\text{No}$	116.2	50.1	$\leq 6.8 \text{ nb}$

Table 5.2: Summary of experimental cold fusion production cross sections. Cross sections without errors are upper limit cross sections only.

Reaction	Energy (MeV)	E* Energy (MeV)	Cross Section
208 Pb(48 Ca,1n) 255 No	204.0	12.2	0.3 ^{+0.6} _{-0.2} nb 2.3 ^{+1.3} _{-1.0} nb
208 Pb $(^{48}$ Ca,1n $)^{255}$ No	206.5	14.2	$2.3^{+1.3}_{-1.0}$ nb
208 Pb(48 Ca,1n) 255 No	207.8	15.2	$9 \pm 3 \text{ nb}$
208 Pb(48 Ca,1n) 255 No	208.8	16.1	100 ⁺⁵⁰ ₋₄₀ nb
²⁰⁸ Pb(⁴⁸ Ca,1n) ²⁵⁵ No	209.0	16.3	$33 \pm 9 \text{ nb}$
²⁰⁸ Pb(⁴⁸ Ca,2n) ²⁵⁴ No	206.5	14.2	$0.6^{+0.8}_{-0.6}$ nb
208 Pb(48 Ca,2n) 254 No	207.8	15.2	1.7 ^{+1.5} _{-1.3} nb
²⁰⁸ Pb(⁴⁸ Ca,2n) ²⁵⁴ No	208.8	16.1	$40 \pm 20 \text{ nb}$
208 Pb(48 Ca,2n) 254 No	209.0	16.3	$58 \pm 15 \text{ nb}$
²⁰⁸ Pb(⁴⁸ Ca,2n) ²⁵⁴ No	211.3	18.1	$900 \pm 300 \text{ nb}$
208 Pb(48 Ca,2n) 254 No	211.5	18.3	$530 \pm 130 \text{ nb}$
208 Pb(48 Ca,2n) 254 No	213.8	20.2	3500 ⁺¹⁰⁰⁰ -900 nb
208 Pb(48 Ca,2n) 254 No	215.3	21.4	$1900 \pm 500 \text{ nb}$
²⁰⁸ Pb(⁴⁸ Ca,2n) ²⁵⁴ No	216.3	22.2	$2300 \pm 600 \text{ nb}$
²⁰⁸ Pb(⁴⁸ Ca,2n) ²⁵⁴ No	218.8	24.2	$1700 \pm 500 \text{ nb}$
²⁰⁸ Pb(⁴⁸ Ca,2n) ²⁵⁴ No	221.4	26.3	$530 \pm 160 \text{ nb}$
208 Pb(48 Ca,3n) 253 No	230.6	33.8	53 ± 14 nb
²⁰⁸ Pb(⁴⁸ Ca,4n) ²⁵² No	230.6	33.8	$1.8^{+1.1}_{-0.8} \; \mathrm{nb}$
208ph (50T; 1 =)257p.c	226.0	12.0	< 0.00¢ 1
208 Pb(50 Ti,1n) 257 Rf	226.0	12.9	$\leq 0.086 \text{ nb}$
²⁰⁸ Pb(⁵⁰ Ti,1n) ²⁵⁷ Rf	231.0	16.9	$16 \pm 4 \text{ nb}$
208 Pb(50 Ti,1n) 257 Rf	233.1	18.6	$16 \pm 5 \text{ nb}$
²⁰⁸ Pb(⁵⁰ Ti,1n) ²⁵⁷ Rf	236.1	21.0	$10 \pm 3 \text{ nb}$
²⁰⁸ Pb(⁵⁰ Ti,2n) ²⁵⁶ Rf	231.0	16.9	$0.6^{+0.5}_{-0.3} \text{ nb} 1.1^{+0.8}_{-0.5} \text{ nb}$
²⁰⁸ Pb(⁵⁰ Ti,2n) ²⁵⁶ Rf	233.1	18.6	
208 Pb(50 Ti,2n) 256 Rf	236.1	21.0	$6.7 \pm 1.8 \text{ nb}$
208 Pb(51 V,2n) 257 Db	250.0	24.9	$1.4^{+0.7}_{-0.6}~{\rm nb}$
²⁰⁹ Bi(⁵⁰ Ti,1n) ²⁵⁸ Db	231.2	15.0	≤ 0.33 nb
209 Bi $(^{50}$ Ti,1n $)^{258}$ Db	233.5	16.9	$3.1^{+1.5}$ _{-1.3} nb
²⁰⁹ Bi(⁵⁰ Ti,2n) ²⁵⁷ Db	233.5	16.9	$\leq 0.25 \text{ nb}$
209 Bi $(^{51}$ V,2n $)^{258}$ Sg	245.4	20.3	14 ⁺³² ₋₁₂ pb
$209 \text{ Pi}(51 \text{ V. 2n})^{258} \text{ Ca}$	243.4 248.4	20.3 22.7	50^{+30}_{-20} pb
209 Bi $(^{51}$ V,2n $)^{258}$ Sg 209 Bi $(^{51}$ V,2n $)^{258}$ Sg	246.4 251.4	25.2	26 +26 ₋₁₅ pb
DI(V,2II) Sg	231.4	<i>43.4</i>	20 -15 pu

Gaussian curves were used to develop a generalized set of systematics and to observe similarities and differences between experimental excitation functions observed at different institutions as well as obtain information about the shape of cold and hot fusion excitation functions. Gaussian curves were used as the shape of excitation functions near their centroid are Gaussian in shape. Using the computer program MathCAD, a simple Gaussian equation, Equation 5.1, was used to fit the experimental data from GSI for two reactions, 208 Pb + 50 Ti and 209 Bi + 50 Ti. These reactions were chosen as the cross sections for the 1n- and 2n-exit channels have been determined at more than 5 different excitation energies [He β 1997, He β 2001]. The Gaussian function used had the form:

$$g(e) = \frac{g_o}{\sqrt{2\pi\sigma}} \exp\left[\frac{-(e-m)^2}{2\sigma^2}\right]$$
 (5.1)

where " g_0 " is the area of the Gaussian function, "m" is the centroid of the Gaussian function, " σ " is the standard deviation and the energy, "e", is the excitation energy of the function.

The fit to the experimental data from GSI was performed by altering the Gaussian curve by adjusting the width (σ) , centroid (m) and magnitude (g_o) until the curve agreed with the experimental data points. Figure 5.1 shows the results of the adjusting the Gaussian curves to the GSI experimental data. Table 5.3 contains the results of the fits to the experimental data.

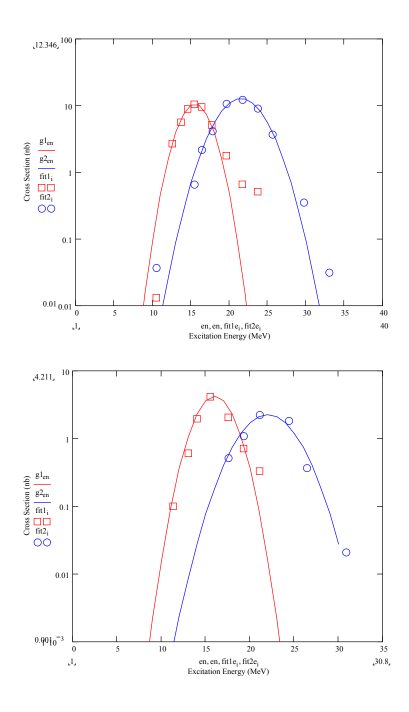


Figure 5.1: Gaussian curves (solid lines) and the experimental data for the $^{208}Pb(^{50}Ti,1n)^{257}Rf$ (top 0), $^{208}Pb(^{50}Ti,2n)^{256}Rf$ (top #), $^{209}Bi(^{50}Ti,1n)^{258}Db$ (bottom 0) and $^{209}Bi(^{50}Ti,2n)^{257}Db$ (bottom #) reactions [He β 1997, He β 2001].

Table 5.3: Results of fits of the Gaussian curves to the experimental data from GSI for the $^{208}Pb(^{50}Ti,xn)^{258-x}Rf$ and $^{209}Bi(^{50}Ti,xn)^{259-x}Db$ reactions. The standard deviation and centroid are labeled " σ " and "m" and the ratio of the areas is " g_o2/g_o1 ".

Projectile	Target	σΙ	m1 (MeV)	σ2	m2 (MeV)	g _o 2 / g _o 1
²⁰⁸ Pb	⁵⁰ Ti	1.8	15.5	2.7	21.5	1.8
²⁰⁹ Bi	⁵⁰ Ti	1.8	16.0	2.7	22.0	0.79

Several important things can be seen from these Gaussian curves. The same standard deviations were used for the 1n- and 2n-exit channel excitation functions for the $^{208}\text{Pb} + ^{50}\text{Ti}$ and $^{209}\text{Bi} + ^{50}\text{Ti}$ reactions. The standard deviations from these fits correspond to FWHM for these excitation functions of 4.2 MeV for the 1n-exit channel and 6.4 MeV for the 2n-exit channel. A difference of 6 MeV in excitation energy for the two centroids was seen for the two experiments. Different areas were used to fit the different excitation functions and therefore different initial area ratios were used. Deviations were seen on the high excitation energy side of the excitation function as the decrease in the excitation function due to fission survivability is much less than the decrease in the excitation function due to the effects of the Coulomb barrier on the low excitation energy side of the excitation function.

The experimental results for the ²⁰⁸Pb(⁵⁰Ti,xn)^{258-x}Rf and ²⁰⁹Bi(⁵⁰Ti,xn)^{259-x}Db reactions from this thesis as well as the other experimental results were then fit to Gaussian curves using the similar centroids, centroid differences, and ratios of 2n to 1n initial areas. The actual initial areas were changed, but the ratios were kept similar. The centroids were altered to make the Gaussian curves appear to model the experimental data. Figure 5.2 shows the Gaussian curve results for the ²⁰⁸Pb(⁵⁰Ti,xn)^{258-x}Rf and ²⁰⁹Bi(⁵⁰Ti,xn)^{259-x}Db experimental results of this thesis. Figure 5.3 illustrates the results of the Gaussian curves to the ²⁰⁸Pb(⁴⁸Ca,xn)^{256-x}No and ²⁰⁹Bi(⁵¹V,xn)^{260-x}Sg reactions. Finally, Figure 5.4 shows the results of the ²⁰⁸Pb(⁵¹V,xn)^{259-x}Db reaction. Table 5.4 summarizes the results of the Gaussian curve fits to the experimental data.

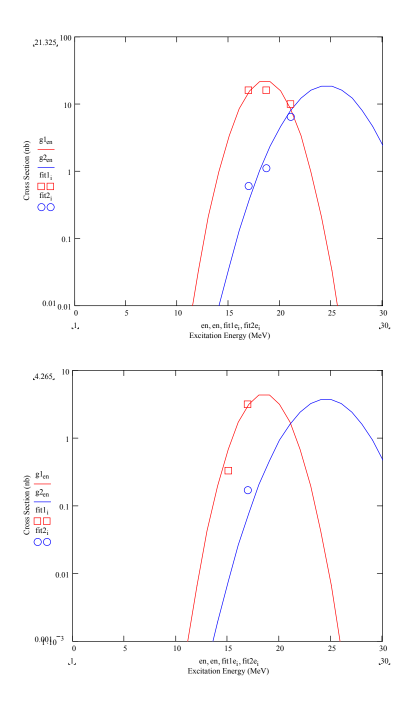


Figure 5.2: Gaussian curves (solid lines) and the experimental data from this thesis for the $^{208}\text{Pb}(^{50}\text{Ti},1n)^{257}\text{Rf}$ (top 0), $^{208}\text{Pb}(^{50}\text{Ti},2n)^{256}\text{Rf}$ (top #), $^{209}\text{Bi}(^{50}\text{Ti},1n)^{258}\text{Db}$ (bottom 0) and $^{209}\text{Bi}(^{50}\text{Ti},2n)^{257}\text{Db}$ (bottom #) reactions. The only data point in the $^{209}\text{Bi}(^{50}\text{Ti},2n)^{257}\text{Db}$ reaction is an upper limit cross section only.

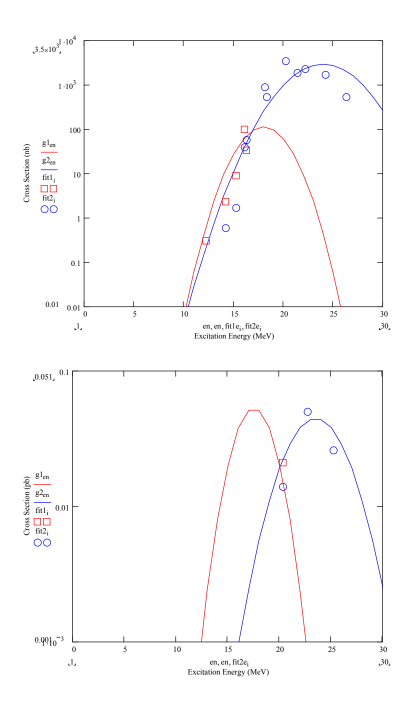


Figure 5.3: Gaussian curves (solid lines) and the experimental data from this thesis for the $^{208}Pb(^{48}Ca,1n)^{255}No~(top~0),~^{208}Pb(^{48}Ca,2n)^{254}No~(top~\#),~^{209}Bi(^{51}V,1n)^{259}Sg~(bottom~0)$ and $^{209}Bi(^{51}V,2n)^{258}Sg~(bottom~\#)$ reactions. The only data point in the $^{209}Bi(^{51}V,1n)^{259}Sg~(bottom~\#)$ reaction is an upper limit cross section only.

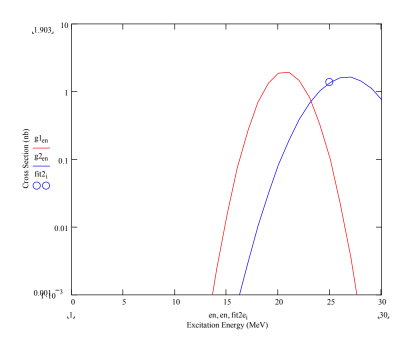


Figure 5.4: Gaussian curves (solid lines) and the experimental data from this thesis for the $^{208}\text{Pb}(^{51}\text{V},2\text{n})^{257}\text{Db}$ (#) reaction.

Table 5.4: Results of Gaussian fits to the experimental data from this thesis for the $^{208}Pb(^{50}Ti,xn)^{258-x}Rf,\,^{209}Bi(^{50}Ti,xn)^{259-x}Db,\,^{208}Pb(^{48}Ca,xn)^{256-x}No,\,^{209}Bi(^{51}V,xn)^{260-x}Sg,$ and $^{208}Pb(^{51}V,xn)^{259-x}Db$ reactions. The standard deviation and centroid are labeled " σ " and "m" and the ratio of the areas is " g_o2/g_o1 ".

Projectile	Target	σ1	m1 (MeV)	σ2	x2 (MeV)	g _o 2 / g _o 1
²⁰⁸ Pb	⁵⁰ Ti	1.8	18.5	2.7	24.5	1.25
²⁰⁹ Bi	⁵⁰ Ti	1.8	18.5	2.7	24.5	1.25
²⁰⁸ Pb	⁴⁸ Ca	1.8	18.0	2.7	24.0	40
²⁰⁹ Bi	^{51}V	1.8	17.5	2.7	23.5	1.25
²⁰⁸ Pb	^{51}V	1.8	18.0	2.7	24.0	1.25

In Figure 5.2, the Gaussian fits to the experimental data were quite good. The only observable discrepancy was the movement of the centroid to higher excitation energies on the order of 2.5 to 3 MeV. Otherwise, keeping the difference in centroid energy between the 1n- and 2n-exit channels constant and the standard deviations constant, the Gaussian fits seemed to predict and model the experimental results. The one point in the ²⁰⁹Bi(⁵⁰Ti,xn)^{259-x}Db reaction for the 2n-exit channel was an upper limit showing that it was probable that no events were seen. The Gaussian fit to the ²⁰⁸Pb(⁴⁸Ca,xn)^{256-x}No experimental data was not as good as other Gaussian fits were to the other experimental data. It was possible that the 2n-exit channel was so large that the In-exit channel was masked. Adding to this problem were the similar decay energies and half-lives of ²⁵⁵No and ²⁵⁴No, the 1n- and 2n-exit channel products. An increase in the ratio $g_0(2n)/g_0(1n)$ was also seen for this reaction which was unlike the other cold fusion reactions. Better fits were obtained with a smaller 2n standard deviation and a larger area ratio, but these fitting parameters were dissimilar to the rest of the cold fusion fit parameters. The fit to the ²⁰⁹Bi(⁵¹V,xn)^{260-x}Sg was performed using the same standard deviations as the other experiments, and the fact that no ²⁵⁹Sg was seen in the experiment at the lowest experimental excitation energy, a guess in terms of the initial area ratio was taken to reflect what was seen experimentally. The parameters for the fit to the experimental ²⁰⁸Pb(⁵¹V,xn)^{259-x}Db data point is speculative as only one experimental point was obtained. With only one point, an infinite number of Gaussian fits are possible, and the one proposed is only speculative.

From the Gaussian fit results for the cold fusion reactions, it was seen that the widths of these excitation functions, the excitation energies of the centroids, and the

difference in excitation energy between the centroids were all consistent. A similar attempt was made to fit the experimental results of the hot fusion reactions using the fits to the cold fusion experiments as an initial guess.

Knowing that the excitation energy at the Coulomb barrier is higher for the hot fusion reactions than the cold fusion reactions, and the fact that the 4n-exit channel is favored over the 3n-exit channel led to an initial guess for the parameters used to fit the experimental results for the hot fusion reactions from this thesis. The standard deviations were slightly higher than those observed for the cold fusion reactions. Understandably the centroids were located at higher excitation energies, and the difference between the centroids for the 3n- and 4n-exit channels was larger than the differences seen in the cold fusion reactions. The $g_0(4n)/g_0(3n)$ ratio was also larger than the $g_0(2n)/g_0(1n)$ ratio observed in the cold fusion reactions. The results to the Gaussian fits can be seen in Figures 5.5 and 5.6, with the summary of the Gaussian fit parameters located in Table 5.5.

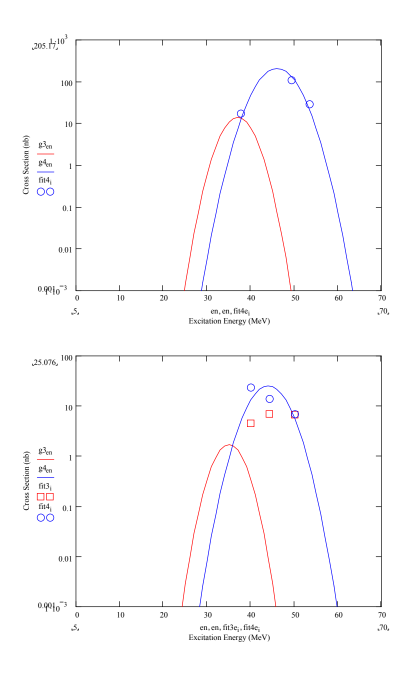


Figure 5.5: Gaussian curves (solid lines) and the experimental data from this thesis for the 238 U(18 O,4n) 252 Fm (top #), 238 U(22 Ne,3n) 257 No (bottom 0) and 238 U(22 Ne,4n) 256 No (bottom #) reactions. The data points in the 238 U(22 Ne,3n) 257 No reaction are upper limit cross sections only.

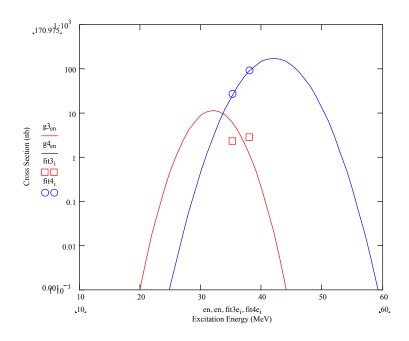


Figure 5.6: Gaussian curves (solid lines) and the experimental data from this thesis for the $^{248}\text{Cm}(^{15}\text{N},3\text{n})^{260}\text{Lr}$ (0) and $^{248}\text{Cm}(^{15}\text{N},4\text{n})^{259}\text{Lr}$ (#) reactions.

Table 5.5: Results of Gaussian fits to the experimental data from this thesis for the $^{238}U(^{18}O,xn)^{256\text{-x}}Fm,\,^{238}U(^{22}Ne,xn)^{260\text{-x}}No,$ and $^{248}Cm(^{15}N,xn)^{263\text{-x}}Lr$ reactions. The standard deviation and centroid are labeled " σ " and "m" and the ratio of the initial areas is " g_o4/g_o3 ".

Projectile	Target	σ3	m3 (MeV)	σ4	m4 (MeV)	g_o4/g_o3
²³⁸ U	¹⁸ O	2.8	37	3.5	46	18.75
^{238}U	²² Ne	2.8	37	3.5	46	18.75
²⁴⁸ Cm	¹⁵ N	2.8	32	3.5	42	18.75

These same Gaussian curves were fit to the data from Donets et al. [Don1966] for the ²³⁸U(¹⁸O,xn)^{256-x}Fm and ²³⁸U(²²Ne,xn)^{260-x}No reactions and Eskola et al. [Esk1971] for the ²⁴⁸Cm(¹⁵N,xn)^{263-x}Lr reaction. The same standard deviations, similar centroids and similar centroid differences for the 3n- and 4n-exit channel excitation functions were found. The standard deviations from Table 5.5 correspond to FWHM values of 6.6 and 8.2 MeV for the 3n- and 4n-exit channel excitation functions. The results in Figure 5.5 are self-explanatory. The parameters for the Gaussian fits match the data very well and in the case of the ²³⁸U(²²Ne,3n)²⁵⁷No illustrates the fact that no ²⁵⁷No should have been seen in the reactions as the cross sections for the 3n-exit channel in that reaction are extremely small at those energies. The Gaussian fit to the ²⁴⁸Cm(¹⁵N,xn)^{263-x}Lr experimental data is also good. The error bars on the cross sections for the 3n-exit channel are in line with the Gaussian fit.

Gaussian curves can be used to help model the shape of excitation functions for various hot and cold fusion reactions around their centroids. Information was obtained from these curves (Tables 5.4 and 5.5) that shows a possible systematic link between the 1n-, 2n-exit channels from the cold fusion reactions and 3n- and 4n- exit channels from the hot fusion reactions. Additional data points for the cold and hot fusion excitation functions mentioned in this thesis would help in the development of modeling the excitation functions with a Gaussian curve and in the understanding of the trends in the shapes of cold and hot fusion excitation functions.

5.2 Exit channel systematics

Cross sections produced in various reactions can be grouped according to projectile isospin (isospin = (number of neutrons – number of protons) \div 2) and then plotted as a function of the maximum experimental cross section of an excitation function for a given exit channel versus the atomic number of the isotope produced. These cross sections are grouped by projectile isospin as the isospins of the targets and resulting compound nuclei are nearly equal. Therefore, the only real difference in these reactions is the projectile isospin. In most of these plots, a decrease in production cross section is seen with an increase in the atomic number of the product. Comparing experimental cross sections to the trend seen systematically can show whether the various cross section values are consistent or not.

The experimental results for the 1n-exit channel reactions were compared to other 1n-exit channel production cross sections in Figure 5.7. For the 1n-exit channel it appears that all of the values were similar to what is seen systematically.

In a similar manner the experimental results for the 2n-exit channel were compared to similar reactions with isospins of 2.5, 3 and 4. The 2n-exit channel experimental results are also consistent with the systematics presented here. The energies studied experimentally included the maxima of the 2n-exit channel excitation functions and therefore give good agreement with the systematics. In one of the experiments only an upper limit value was obtained for the 2n-exit channel and therefore the corresponding data point in Figure 5.8 is not in good agreement with the systematics. Figure 5.8 shows the results of the comparison.

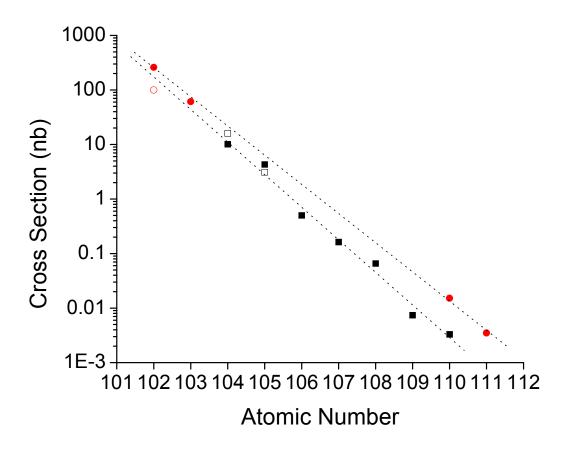


Figure 5.7: 1n-exit channel systematics (solid symbols +) compared with experimental results (open symbols #). Square symbols (+) represent an isospin of 3 and circle symbols (!) an isospin of 4 [Gäg1989, He β 1997, He β 2001, Mün1985, Mün1989, Hof1998, Hof1997, Hof2001, Hof1995a]. The dotted lines are to guide the eye along cross sections from the same isospin.

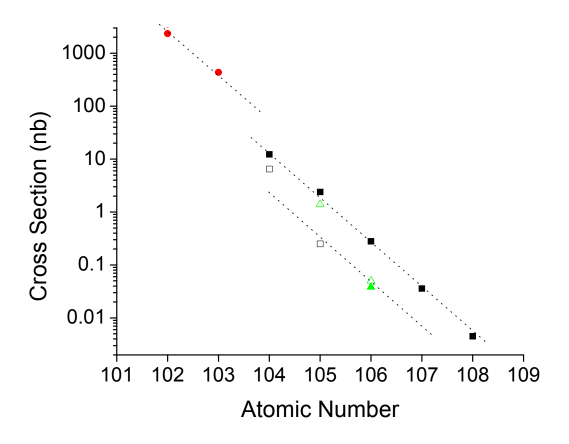


Figure 5.8: 2n-exit channel systematics (solid symbols +) compared with experimental results (open symbols 0). Triangle symbols (:) represent an isospin of 2.5, square symbols (+) an isospin of 3, and circle symbols (!) an isospin of 4 [Gäg1989, Heβ1997, Heβ2001, Mün1985, Mün1989, Hof1998]. The dotted lines are to guide the eye along cross sections from the same isospin.

Finally, the experimental 4n-exit channel data were compared with systematics for the 4n-exit channel. No 3n-exit channel comparison was done as there is not enough experimental data present to make any systematic study. There are quite a few ways to produce elements heavier than einsteinium in hot fusion reactions as there is a variety of target and projectiles to work with. With the hot fusion reactions, a specific projectile was chosen to do the comparison with experimental data. A significant amount of experimental work has been done with ²²Ne projectiles and therefore these reactions were chosen to do the comparison for the 4n-exit channel. Figure 5.9 shows the comparison of the ²²Ne experimental work of this thesis (open symbol) compared with other ²²Ne experimental work making isotopes of the elements from fermium to bohrium (solid symbols). The experimental results of this thesis are consistent with the systematic trends.

The comparison of the experimental production cross sections with the systematics of the 1n-, 2n- and 4n-exit channel seems rather straight forward. In a majority of the cases, the data seem to correspond well with what has been seen previously and in a few cases, larger production cross sections might have been seen for a particular reaction if the excitation function had been studied in greater detail.

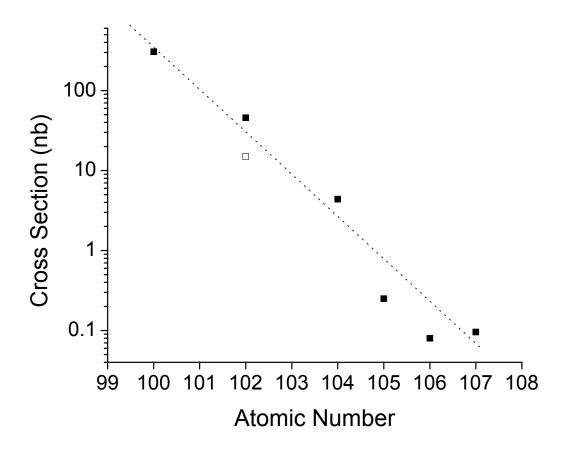


Figure 5.9 : 4n-exit channel systematics (solid symbols +) compared with experimental results of this thesis (open symbols 0) for hot fusion reactions with ²²Ne projectiles [Tür1999, Hof1998, Laz2000, Lan1988, Laz1994, Wil2000]. The dotted line is present to guide the eye.

5.3 HIVAP cross sections

The comparisons between the experimental cross sections of this thesis and the predictions of the HIVAP code with the Reisdorf and Schädel parameters were presented in Section 4. The Reisdorf and Schädel input parameters were satisfactory in predicting the cross sections for the 3n- and 4n-exit channel excitation functions from the hot fusion reactions, but were not statisfactory at predicting the 1n- and 2n-exit channel excitation functions for the cold fusion reactions.

The Reisdorf and Schädel input parameters for HIVAP were altered in the attempt to model the experimental results more accurately. These alterations were called the Patin parameters. A full list of the input parameters as they appear in the input file can be found in Appendix B. The aim was to alter as few of the parameters as possible. Only six of the Reisdorf and Schädel input parameters were changed: LEVELPAR, AF/AN, V0, Q2, SIGR0, and CUTOFF. LEVELPAR is the value of the radius parameter that is used when calculating the level density ratios for the de-excitation steps. LEVELPAR was altered to 1.16, the value used in the calculations of Töke and Swiatecki [Tök1981]. AF/AN is an integer that determines what designates the calculation to be used. In the Reisdorf and Schädel parameter set, AF/AN equals 1 meaning the level density ratio was calculated using Reisdorf [Rei1981]. The Patin parameters used an AF/AN value equal to 2 to designate using the level density calculations of Töke and Swiatecki [Tök1981]. The Töke/Swiatecki calculation better reflects what occurs experimentally in the cold fusion reactions. V0 equals the nuclear well potential for the target atom which was changed to reflect a smaller potential well depth of about 50 MeV rather than 70 MeV used in the Reisdorf and Schädel parameters. Q2 is the quadrupole moment of the target

atom in units of fm². A Q2 value of 0 was used to reflect that in the cold fusion reactions, the lead and bismuth targets are essentially spherical. The SIGR0 value is the percent fluctuation in R0 used to calculate the fusion barrier in the entrance channel used for determining the fusion cross section. CUTOFF is the integration limit in units of SIGR0. SIGR0 and CUTOFF influence the entrance channel fusion barrier and were altered depending on the specific reactions to reflect the wide differences between the predictions of the HIVAP code using the Reisdorf and Schädel parameters and the experimental data. The values of SIGR0 varied between 1.7 and 4.0 percent. The CUTOFF value was left at 4.0 or 5.0.

The input parameter values LEVELPAR, AF/AN, V0 and Q2 were left at their altered values when comparing the predictions of the Patin parameters to the experimental data. The values of SIGR0 and CUTOFF were varied to reflect the changes in the projectile and targets used in the reactions. The largest effect in the predictions of the HIVAP code came by as a result of the alterations to the entrance channel values of SIGR0 and CUTOFF. A small barrier fluctuation value was chosen for the ²⁰⁸Pb(⁴⁸Ca,xn) reaction because of the spherical character of both the projectile and target. As the projectiles moved away from the sphericity of ⁴⁸Ca and the target was changed to ²⁰⁹Bi, the barrier fluctuation value was increased. Figures 5.10, 5.11, 5.12 and 5.13 show a comparison between the predictions of HIVAP using the Patin parameters, the experimental data of this thesis, and experimental results obtained elsewhere.

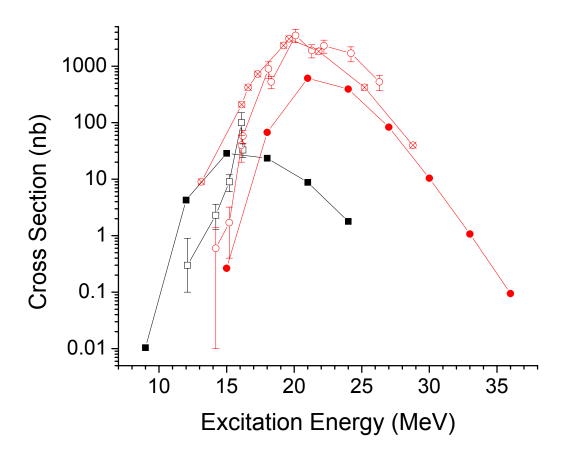


Figure 5.10: Predictions of the HIVAP code using the Patin parameters (solid symbols .,!), the experimental production cross sections (open symbols 0,#), and previous experimental results [Gäg1989] (open symbols with an 4) for the ²⁰⁸Pb(⁴⁸Ca,1n)²⁵⁵No (squares) and ²⁰⁸Pb(⁴⁸Ca,2n)²⁵⁴No (circles) reactions. A SIGR0 value of 1.7 and a CUTOFF value of 5.0 were used in the input parameter set.

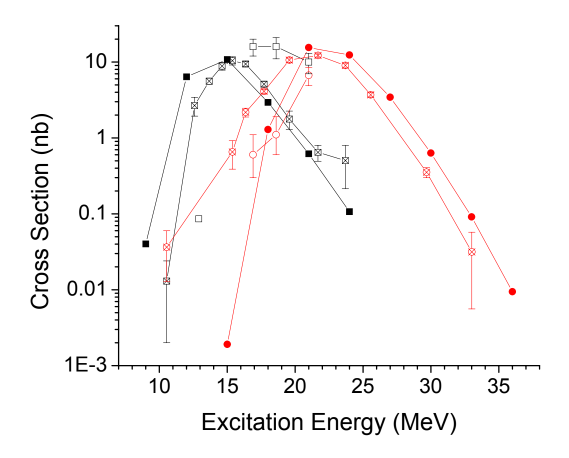


Figure 5.11: Predictions of the HIVAP code using the Patin parameters (solid symbols .,!), the experimental production cross sections (open symbols 0,#), and previous experimental results [He β 1997] (open symbols with an 4) for the 208 Pb(50 Ti,1n) 257 Rf (squares) and 208 Pb(50 Ti,2n) 256 Rf (circles) reactions. A SIGR0 value of 2.6 and a CUTOFF value of 5.0 were used in the input parameter set.

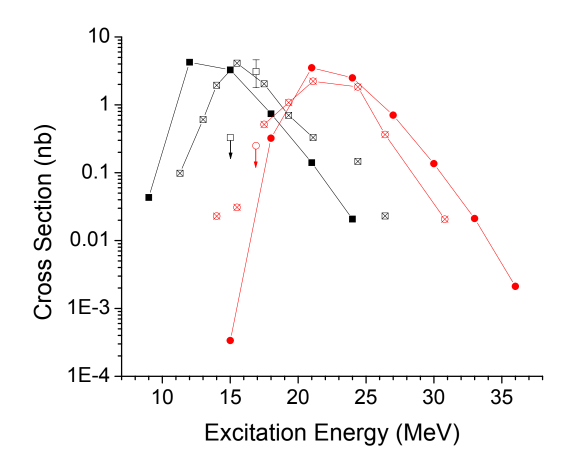


Figure 5.12: Predictions of the HIVAP code using the Patin parameters (solid symbols .,!), the experimental production cross sections (open symbols 0,#), and previous experimental results [He β 2001a] (open symbols with an 4) for the 209 Bi(50 Ti,1n) 258 Db (squares) and 209 Bi(50 Ti,2n) 257 Db (circles) reactions. A SIGR0 value of 2.9 and a CUTOFF value of 5.0 were used in the input parameter set.

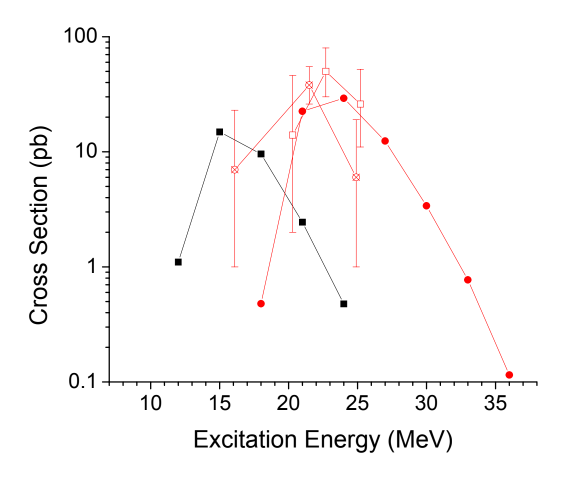


Figure 5.13: Predictions of the HIVAP code using the Patin parameters (solid symbols .,!), the experimental production cross sections (open symbols 0,#), and previous experimental results [He β 2001b] (open symbols with an 4) for the $^{209}\text{Bi}(^{51}\text{V},1n)^{259}\text{Sg}$ (squares) and $^{209}\text{Bi}(^{51}\text{V},2n)^{258}\text{Sg}$ (circles) reactions. A SIGR0 value of 4.0 and a CUTOFF value of 4.0 were used in the input parameter set.

The alterations to the input parameters seem to have a dramatic effect on the predictions of the HIVAP code in comparison to the predictions of the HIVAP code using the Reisdorf and Schädel parameters. The Patin parameters more accurately reflect what is occurring in the cold fusion reactions. The Reisdorf and Schädel parameter set was tailored for hot fusion reactions with actinide targets and accurately predicts those particular production cross sections. Barrier fluctuations between different targets and projectiles aren't as important in the hot fusion reactions as they don't change as drastically from target to target, and projectile to projectile. In addition, the hot fusion reactions that were modeled by Reisdorf and Schädel occurred at energies well above the barrier where these fluctuations do not have as much influence on the production cross section. The Patin parameters are more accurate as a result of tailoring the input parameters to these specific cold fusion reactions. The differences in the percentage of fusion barrier fluctuation (SIGR0) were discovered through comparison of the HIVAP predictions using the altered parameters and the experimental data of this thesis. Very little barrier fluctuation was require to model the ²⁰⁸Pb(⁴⁸Ca,xn) reaction whereas increasingly larger barrier fluctuation values were required for the ²⁰⁸Pb(⁵⁰Ti,xn), ²⁰⁹Bi(⁵⁰Ti,xn), and ²⁰⁹Bi(⁵¹V,xn) reactions. The following trend of increasing barrier fluctuation value SIGR0 was observed: $^{208}Pb(^{50}Ti,2n)^{256}Rf < ^{209}Bi(^{50}Ti,2n)^{257}Db < ^{50}Ti,2n$ 208 Pb(51 V,2n) 257 Db < 209 Bi(51 V,2n) 258 Sg. An effective input parameter set has been found to predict the production cross sections of cold fusion reactions by allowing for small variations in the SIGR0 and CUTOFF parameters.

5.4 Odd particle effects

Using two different targets, ²⁰⁸Pb and ²⁰⁹Bi, and two different projectiles, ⁵⁰Ti and ⁵¹V, four different reactions can occur. In this case, two of the reactions, ²⁰⁸Pb(⁵¹V,xn)^{259-x}Db and ²⁰⁹Bi(⁵⁰Ti,xn)^{259-x}Db produce the same compound nucleus. By studying these four different reactions, insights can be obtained into the role of the odd proton in the target (²⁰⁹Bi + ⁵⁰Ti), or in the projectile (²⁰⁸Pb + ⁵¹V), or in both (²⁰⁹Bi + ⁵¹V) when compared to the even proton projectile and target system (²⁰⁸Pb + ⁵⁰Ti). Using the experimental results of this thesis and the predictions of HIVAP using the altered parameters from Section 5.3, it appears that the production cross sections can vary by a factor of 300 depending on where the odd proton is located.

As a standard, the cross sections for the 2n-exit channel were used from each of the four reactions. The maximum in the ²⁰⁸Pb(⁵⁰Ti,2n)²⁵⁶Rf reaction had a cross section of about 10 nb. When the odd proton was located in the projectile as in the ²⁰⁸Pb(⁵¹V,2n)²⁵⁷Db reaction, the maximum cross section was about 2 nb. If the odd proton was located in the target as in the ²⁰⁹Bi(⁵⁰Ti,2n)²⁵⁷Db reaction, the maximum cross section was about 3 nb. In this case, a larger cross section was obtained in the reaction of the even Z projectile with the odd Z target. When both the target and projectile had an odd proton as in the ²⁰⁹Bi(⁵¹V,2n)²⁵⁸Sg reaction, the maximum cross section decreased by a factor of more than 200 to 50 pb. These results were seen in both the experimental data as well as the predictions of the HIVAP code with both the Reisdorf and Schädel parameters and the Patin input parameters. These results were also seen in the predictions of the SPIT code. The specific ratios between the maxima were different, but the trend for the maximum of the 2n-exit channel cross sections was always the same:

²⁰⁸Pb(⁵⁰Ti,2n)²⁵⁶Rf > ²⁰⁹Bi(⁵⁰Ti,2n)²⁵⁷Db > ²⁰⁸Pb(⁵¹V,2n)²⁵⁷Db >> ²⁰⁹Bi(⁵¹V,2n)²⁵⁸Sg.
⁵²Cr, a projectile with the same number of neutrons as ⁵⁰Ti and ⁵¹V, would according to these systematics have a smaller maximum 2n-exit channel cross section from the
²⁰⁸Pb(⁵²Cr,2n)²⁵⁸Sg and ²⁰⁹Bi(⁵²Cr,2n)²⁵⁹Bh reactions than either the ²⁰⁸Pb(⁵¹V,2n)²⁵⁷Db and ²⁰⁹Bi(⁵¹V,2n)²⁵⁸Sg reactions. Predictions using the HIVAP code with the Patin input parameters also predict a smaller 2n-exit channel maximum cross section. The SPIT code predicts similar 2n-exit channel maximum cross sections for all three of the reactions: ²⁰⁹Bi(⁵¹V,2n)²⁵⁸Sg, ²⁰⁸Pb(⁵²Cr,2n)²⁵⁸Sg, and ²⁰⁹Bi(⁵²Cr,2n)²⁵⁹Bh. It would be worthwhile to obtain experimental evidence to back up these predictions, especially if the cross sections are in the picobarn to tens of picobarns range.

The significant difference in the maximum cross section for the ²⁰⁹Bi(⁵¹V,2n)²⁵⁸Sg reaction versus the ²⁰⁸Pb(⁵⁰Ti,2n)²⁵⁶Rf, ²⁰⁹Bi(⁵⁰Ti,2n)²⁵⁷Db, and ²⁰⁸Pb(⁵¹V,2n)²⁵⁷Db reactions cannot be explained by large differences in Q-value, neutron separation energy differences, or by shell effects. Another explanation must exist. One possible explanation was obtained using the calculations of Blocki and Swiatecki [Blo1982, Swi2002]. For each reaction, the deformation energy of the optimized neck configuration of the combined system can be plotted versus the length between the projectile and target centers minus their radii. These plots can be seen in Figure 5.14. The curve for a particular reaction represents the additional barrier which must be overcome after the target and projectile reach a touching configuration. The difference between the deformation energy at the maximum of the curve (saddle point) and the deformation energy at the injection point (0-3 fm for all of these reactions) can then be placed in an exponential function to arrive at the probability of overcoming the barrier to compound

nucleus formation. These probabilities are plotted in Figure 5.15 for various injection point distances.

It is seen from these calculations that the difference in energy between the maximum of the curve and the injection point increases as Z increases, which leads to the decreased probability of forming the compound nucleus. The probability of compound nucleus formation also decreases as the distance of the injection point increases. If the injection point is held at a constant length, the target and projectile system have a larger barrier to surmount to reach the complete fusion of the compound nucleus, and will more likely slide down the barrier to fission. This is somewhat similar to what is seen experimentally. This explanation is only an attempt at explaining the large difference in the 2n-exit channel cross sections seen. It is not a complete explanation, but it does give some insight into what possibly might be occurring in these reactions.

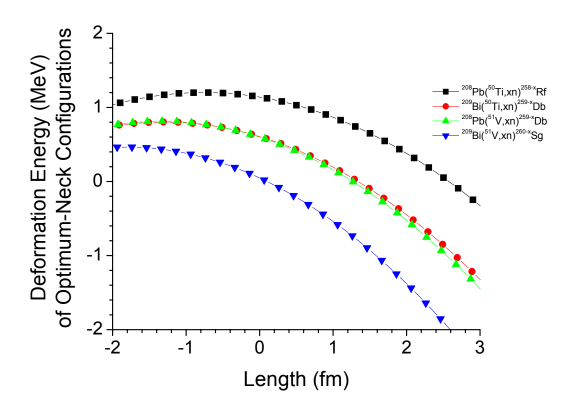


Figure 5.14: Plot of the deformation energy (MeV) of an optimum neck configuration versus length (fm) [Blo1982, Swi2002] for various cold fusion reactions.

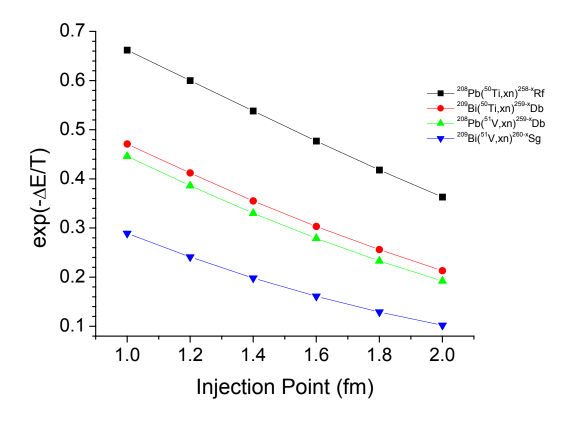


Figure 5.15: Plot of the probability of overcoming the barrier from Figure 5.14 versus the injection point length (fm). ΔE is the difference in deformation energy at the maximum of the curve and the injection point. The temperature T was calculated assuming an excitation energy of 20 MeV and a level density parameter a equal to A/8.5 MeV.

6 Conclusions and future research

6.1 Conclusions

The MG system is an effective system to study the production of isotopes with fairly significant cross sections (> 1 nb). Studies of heavier system become increasingly difficult as the cross sections become smaller. The catcher foil technique is useful when studying long-lived heavy element isotopes but not as effective when studying the short-lived isotopes of the heaviest elements. The BGS is extremely successful for the study of cold fusion reactions. The large difference between the magnetic rigidities of the evaporation residues and transfer products makes separation of cold fusion reaction products highly efficient. The BGS efficiency for hot fusion reactions using lighter beams is significantly lower and therefore reduces the usefulness of the BGS in the study of hot fusion reactions. Overall, as the study of hot fusion reactions continues, new techniques are needed to observe the short half-life neutron rich heavy element activities produced with heavy ion beams and actinide targets. The BGS should continue to be used to study the various effects in cold fusion reactions.

The non-observation of the 3n-exit channel in the hot fusion ²³⁸U target reactions indicates that the 3n-exit channel is more difficult to observe than anticipated. One possible explanation is the sub-Coulomb barrier fusion required for these reactions to be successful. The sub-Coulomb barrier energies coupled with the 3-neutron evaporation steps at which depletion by fusion can occur leaves the 3n-exit channel as a difficult route for the production of the neutron-rich isotopes of the heaviest elements compared to the 4n-exit channel. As the 4n-exit channel is closer to the Coulomb barrier, the fusion cross

section is higher making the 4n-exit channel a more dominant route to heavy element production. Gaussian modeling confirmed that the fusion barrier has a significant role in the depression of the 3n-exit channel versus the 4n-exit channel in hot fusion reactions.

The observation of equally probable 1n- and 2n-exit channels in cold fusion reactions is no surprise. The same sub-Coulomb barrier hindrance observed in the 3n-exit channel in hot fusion reactions is reduced in cold fusion reactions because of a reduced chance for fission because of only one neutron evaporation step. The increase in the fusion cross section at higher energies is canceled out by the second evaporation step for the 2n-exit channel leaving both the 1n- and 2n-exit channels approximately equal. These results are consistent for all of the cold fusion experiments except the $^{208}\text{Pb}(^{48}\text{Ca},\text{xn})^{256-x}\text{No}$ reaction in which the 2n-exit channel is much larger than the 1n-exit channel. Possible explanations include the significant fusion cross section enhancement at lower excitation energies for the doubly magic projectile and target system.

The HIVAP code using the Reisdorf and Schädel parameters was accurate in predicting the hot fusion reaction cross sections. Modifications to the input parameters of the HIVAP code helped accurately predict the cold fusion production cross sections. It was observed that fusion barrier fluctuations could play an important role in the overall production cross section. These effects were not seen in the hot fusion reactions probably because hot fusion reaction targets are deformed and not spherical like the lead and bismuth targets. The HIVAP code has been shown to be an effective tool for predicting the production cross sections for various reaction mechanisms. It is, however, important to use HIVAP only to predict the cross sections of isotopes in regions where experiments

have already been performed, and input parameters have carefully been tailored. The HIVAP code is only effective when used to predict cross sections in a particular region of the Chart of Nuclides.

6.2 Future research

The observation that the 3n-exit channel is more difficult to see as the projectile Z increases could lead to the demise of future hot fusion reactions for the production of neutron-rich nuclei. The MG system should be used to study the ²³⁸U(²²Ne,3n)²⁵⁷No reaction in more detail in order to learn more about 3n-exit channel systematics. The BGS is not suited to the study of this particular experiment, because the efficiency of the BGS is to low to produce any significant results. Heavier beams would lead to better efficiencies in the BGS, but lower cross sections may negate the gain from the larger efficiency. The reactions of ²³⁸U, ²⁴⁴Pu, and ²⁴⁸Cm targets with high intensity ⁴⁸Ca beams may help provide insight into the 3n-exit channel, but with production cross sections in the picobarn range, the possibility of a successful study is slim.

With efficiencies in the BGS around 45%, the continuing study of cold fusion reactions with ²⁰⁸Pb and ²⁰⁹Bi targets is promising. Additional work needs to be done to complete the 1n- and 2n- exit channel excitation functions for the ²⁰⁸Pb(⁵⁰Ti,xn)^{258-x}Rf, ²⁰⁹Bi(⁵⁰Ti,xn)^{259-x}Db, ²⁰⁸Pb(⁵¹V,xn)^{259-x}Db, and ²⁰⁹Bi(⁵¹V,xn)^{260-x}Sg reactions. Once those are complete, detailed information can be accumulated from the shapes of the excitation functions using the Gaussian fits as well as the predictions of the HIVAP code using the Patin parameters. Studies should be continued by comparing those results with additional experiments with ⁵²Cr and ²⁰⁸Pb and ²⁰⁹Bi targets. These reactions would be useful not only for comparison with the already completed reactions with ⁵⁰Ti and ⁵¹V (N=28) projectiles, but also for the study of the neutron-deficient isotopes of the heaviest elements. Once these experiments have been completed, a completely new set of experiments should be performed using the slightly heavier projectiles ⁵⁴Cr, ⁵⁵Mn, and

 56 Fe (N = 30) with the 208 Pb and 209 Bi targets. These reactions would be important as similarities could be drawn between the results of these experiments and the results of the experiments with 50 Ti and 51 V projectiles that have already been studied. These reactions would also extend the study of the neutron-deficient heavy element isotopes.

All of these experiments are exciting and interesting scientifically and have one main goal, to study the different cross sections for reactions producing the heavy elements.

Appendix A: GOOSY data analysis code

```
X$ANAL:@PROCEDURE(P BUFFER, P EVENT)RETURNS(BIN FIXED(31));
      COMMENTS
______
DESIGNED FOR TYPE 10 1 EVENTS/SUBEVENTS WITH SUBCRATE NUMBER USED TO LABEL
   DIFFERENT SUBEVENTS
EACH EVENT HAS AT LEAST ONE SUBEVENT; THE DATE IS FOUND IN THE SUBEVENT
______
     VARIABLE DECLARATIONS
                                   POINTER; /* to current buffer */
POINTER; /* to current event */
DCL P BUFFER
DCL P EVENT
/* DATA ELEMENTS FOR EL, YLT, YHT, YLB, YHB, BL, BH, PT */
                                   POINTER STATIC;
DCL P EL
                POINTER STATIC;
DCL P EH
                                    POINTER STATIC;
DCL P_YLT DCL P_YHT
DCL P YLB
DCL P_YHB
DCL P_BL
DCL P_BH
DCL P PT
DCL P_SEL
DCL P SEH
DCL P SYLT
DCL P_SYHT
DCL P SYLB
DCL P SYHB
DCL P SBL
DCL P_SBH
DCL P SPT
DCL P CEL
DCL P_CEH
DCL P_CYLT
DCL P_CYHT
DCL P CYLB
DCL P_CYHB
DCL P_CBL
DCL P CBH
DCL P CPT
                                     POINTER STATIC;
/\star DATA ELEMENTS FOR CALIBRATION SLOPES AND INTERCEPTS \star/
               POINTER STATIC:
DCL P ELM
                                    POINTER STATIC;
DCL P_ELB
DCL P EHM
                                     POINTER STATIC;
DCL P EHB
                                    POINTER STATIC;
DCL P YLTM
                                POINTER STATIC;
DCL P_YLTB
DCL P_YLBM
                  POINTER STATIC,
POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
                              POINTER STATIC;
POINTER STATIC;
DCL P YLBB
DCL P_YHTM
DCL P_YHTB
DCL P_YHBM
DCL P_YHBB
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/* DATA ELEMENTS FOR CALIBRATED POSITIONS AND ENERGIES */
              POINTER STATIC;
DCL P PKEV
DCL P PMEV
                             POINTER STATIC;
                            POINTER STATIC;
DCL P PLO
                        POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
DCL P_PHI
DCL P YELT
DCL P YEHT
DCL P_YELB
DCL P_YEHB
DCL P EKEV
                             POINTER STATIC;
                          POINTER STATIC;
                          POINTER STATIC;
DCL P_EMEV
DCL P_ELO
                             POINTER STATIC;
DCL P EHI
                             POINTER STATIC;
/* DATA ELEMENTS FOR CORRELATIONS */
DCL P_E_EVR POINTER STATIC;
                            POINTER STATIC;
POINTER STATIC;
DCL P E MOM
DCL P DTEA
                        POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
DCL P_E_DAU
DCL P_E_DAU
DCL P_E_FISS
DCL P_E_EVRF
DCL P_E_PPAC
/* DATA ELEMENTS NAI AND RUTHERFORDS */
                       POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
POINTER STATIC;
DCL P_NAIL POINTER STATIC;
DCL P_NAIH
DCL P_RESLSW
DCL P_RESMSW
DCL P_RWSLSW
DCL P_RWSMSW
DCL P RE
                          POINTER STATIC;
DCL P RW
                           POINTER STATIC;
/* OTHER DATA ELEMENTS */
DCL P_IPAR
                             POINTER STATIC;
DCL P_SLOPE
                            POINTER STATIC;
DCL P_PPAC
DCL P_ERROR
                           POINTER STATIC;
                             POINTER STATIC;
/**** SYSTEM VARIABLE STRUCTURES AND PROCEEDURES *****/
/**** ----- NOT REFERENCED IN DATA BASE ----- ****/
@INCLUDE $MACRO(DCL PROC);
@INCLUDE $MACRO(S$MESS);
@INCLUDE $MACRO($MACRO);
@INCLUDE $MACRO(U$PRTCL);
@INCLUDE $MACRO(SA$VE10 1);
@INCLUDE $MACRO(SA$VES10 1);
@INCLUDE $MACRO(SA$BUFHE);
@INCLUDE $MACRO(U$RANDOM);
/**** SYSTEM VARIABLE STRUCTURES REFERENCED IN DATA BASE ****/
DCL P SECAM
                 POINTER STATIC;
@INCLUDE $MACRO(SA$secam);
P_SA$SECAM = P SECAM;
DCL 1 S_ BASED(P_),
    2 L_LOW BIN FIXED(31),
    2 L\_HIGH BIN FIXED(31),
    2 R (1 REFER(L LOW):1 REFER(L HIGH)) BIN FLOAT(24);
DCL 1 S EL BASED (P EL),
    2 L_EL_LOW BIN FIXED(31),
2 L_EL_HIGH BIN FIXED(31),
    2 R_EL(1 REFER(L_EL_LOW):1 REFER(L_EL_HIGH)) BIN FLOAT(24);
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DCL 1 S EH BASED (P EH),
    2 L_EH_LOW BIN FIXED(31),
2 L_EH_HIGH BIN FIXED(31),
    2 R EH(1 REFER(L EH LOW):1 REFER(L EH HIGH)) BIN FLOAT(24);
DCL 1 S YLT BASED (P YLT),
    2 L_YLT_LOW BIN FIXED(31),
2 L_YLT_HIGH BIN FIXED(31),
                  BIN FIXED(31),
    2 R YLT (1 REFER(L YLT LOW):1 REFER(L YLT HIGH)) BIN FLOAT(24);
DCL 1 S_YHT BASED(P_YHT),
    2 L YHT LOW
                  \overline{\text{BIN FIXED}(31)},
    2 L YHT HIGH BIN FIXED (31),
    2 R_YHT(1 REFER(L_YHT_LOW):1 REFER(L_YHT_HIGH)) BIN FLOAT(24);
DCL 1 S_YLB BASED(P_YLB),
    2 L YLB LOW BIN FIXED(31),
    2 L YLB HIGH BIN FIXED (31),
    2 R YLB(1 REFER(L YLB LOW):1 REFER(L YLB HIGH)) BIN FLOAT(24);
DCL 1 S_YHB BASED(P_YHB),
2 L YHB LOW BIN FIXED(31),
    2 L YHB HIGH BIN FIXED (31),
    2 R YHB (1 REFER (L YHB LOW): 1 REFER (L YHB HIGH)) BIN FLOAT (24);
DCL 1 S BL BASED (P BL),
    2 L BL LOW BIN FIXED (31),
    2 L_BL_HIGH BIN FIXED(31),
    2 R BL(1 REFER(L BL LOW):1 REFER(L BL HIGH)) BIN FLOAT(24);
DCL 1 S BH BASED (P BH),
    2 L BH LOW BIN FIXED (31),
    2 L_BH_HIGH BIN FIXED(31),
    2 R_BH(1 REFER(L_BH_LOW):1 REFER(L_BH_HIGH)) BIN FLOAT(24);
DCL 1 S PT BASED (P PT),
    2 L_PT_LOW BIN FIXED(31),
2 L PT HIGH BIN FIXED(31),
    2 R_PT(1 REFER(L_PT_LOW):1 REFER(L_PT_HIGH)) BIN FLOAT(24);
DCL 1 S CEL BASED (P CEL),
    2 L_CEL_LOW BIN FIXED(31),
2 L_CEL_HIGH BIN FIXED(31),
    2 R CEL(1 REFER(L CEL LOW): 1 REFER(L CEL HIGH)) BIN FLOAT(24);
DCL 1 S_CEH BASED(P_CEH),
    2 L_CEH_LOW
                  BIN FIXED(31),
    2 L CEH HIGH BIN FIXED (31),
    2 R CEH (1 REFER (L CEH LOW): 1 REFER (L CEH HIGH)) BIN FLOAT (24);
DCL 1 S_CYLT BASED(P_CYLT),
    2 L CYLT LOW BIN FIXED (31),
    2 L CYLT HIGH BIN FIXED (31),
    2 R CYLT(1 REFER(L CYLT LOW):1 REFER(L CYLT HIGH)) BIN FLOAT(24);
DCL 1 S CYHT BASED (P CYHT),
    2 L_CYHT_LOW BIN FIXED(31),
    2 L CYHT HIGH BIN FIXED (31),
    2 R CYHT (1 REFER(L CYHT LOW):1 REFER(L CYHT HIGH)) BIN FLOAT(24);
DCL 1 S_CYLB BASED(P_CYLB),
    2 L CYLB LOW BIN FIXED (31),
    2 L CYLB HIGH BIN FIXED (31),
    2 R_CYLB(1 REFER(L_CYLB_LOW):1 REFER(L_CYLB_HIGH)) BIN FLOAT(24);
DCL 1 S CYHB BASED(P_CYHB),
    2 L CYHB LOW BIN FIXED (31),
    2 L_CYHB_HIGH BIN FIXED(31),
    2 R CYHB(1 REFER(L CYHB LOW):1 REFER(L CYHB HIGH)) BIN FLOAT(24);
DCL 1 S CBL BASED (P CBL),
    2 L_CBL_LOW BIN FIXED(31),
    2 L CBL HIGH BIN FIXED (31),
    2 R_CBL(1 REFER(L_CBL_LOW):1 REFER(L_CBL_HIGH)) BIN FLOAT(24);
DCL 1 S CBH BASED (P CBH),
    2 L_CBH_LOW BIN FIXED(31),
2 L_CBH_HIGH BIN FIXED(31),
    2 R CBH (1 REFER(L CBH LOW):1 REFER(L CBH HIGH))
                                                        BIN FLOAT(24);
DCL 1 S CPT BASED (P CPT),
    2 L_CPT_LOW
                  BIN FIXED(31),
    2 L CPT HIGH BIN FIXED (31),
    2 R CPT (1 REFER(L CPT LOW):1 REFER(L CPT HIGH)) BIN FLOAT(24);
DCL 1 S_SEL BASED(P_SEL),
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2 L SEL LOW BIN FIXED(31),
   2 L SEL HIGH BIN FIXED(31),
    2 R SEL (1 REFER(L SEL LOW): 1 REFER(L SEL HIGH)) BIN FLOAT(24);
DCL 1 S SEH BASED (P SEH),
   2 L SEH LOW BIN FIXED(31),
   2 L_SEH_HIGH BIN FIXED(31),
   2 R_SEH(1 REFER(L_SEH_LOW):1 REFER(L_SEH HIGH)) BIN FLOAT(24);
DCL 1 S SYLT BASED (P SYLT),
   2 L_SYLT_LOW BIN FIXED(31),
   2 L SYLT HIGH BIN FIXED (31),
   2 R SYLT (1 REFER(L SYLT LOW): 1 REFER(L SYLT HIGH)) BIN FLOAT (24);
DCL 1 S SYHT BASED (P SYHT),
   2 L_SYHT_LOW BIN FIXED(31),
2 L SYHT HIGH BIN FIXED(31),
   2 R SYHT(1 REFER(L SYHT LOW):1 REFER(L SYHT HIGH)) BIN FLOAT(24);
DCL 1 S_SYLB BASED(P_SYLB),
    2 L SYLB LOW BIN FIXED (31),
   2 L SYLB HIGH BIN FIXED (31),
   2 R SYLB(1 REFER(L SYLB LOW):1 REFER(L SYLB HIGH)) BIN FLOAT(24);
DCL 1 S_SYHB BASED(P SYHB),
   2 L SYHB LOW BIN FIXED (31),
   2 L SYHB HIGH BIN FIXED (31),
   2 R_SYHB(1 REFER(L_SYHB_LOW):1 REFER(L_SYHB_HIGH)) BIN FLOAT(24);
DCL 1 S SBL BASED (P SBL),
   2 L SBL LOW BIN FIXED(31),
   2 L SBL HIGH BIN FIXED(31),
   2 R SBL (1 REFER(L SBL LOW):1 REFER(L SBL HIGH)) BIN FLOAT(24);
DCL 1 S SBH BASED (P SBH),
   2 L SBH LOW BIN FIXED(31),
   2 L SBH HIGH BIN FIXED(31),
    2 R SBH (1 REFER(L SBH LOW): 1 REFER(L SBH HIGH)) BIN FLOAT(24);
DCL 1 S SPT BASED (P CPT),
   2 L SPT LOW BIN FIXED(31),
   2 L_SPT_HIGH BIN FIXED(31),
   2 R_SPT(1 REFER(L_SPT_LOW):1 REFER(L_SPT_HIGH)) BIN FLOAT(24);
DCL 1 S ELM BASED (P ELM),
   2 L_ELM_LOW BIN FIXED(31),
   2 L ELM HIGH BIN FIXED(31),
   2 R ELM(1 REFER(L ELM LOW):1 REFER(L ELM HIGH)) BIN FLOAT(24);
DCL 1 S ELB BASED (P ELB),
   2 L_ELB_LOW BIN FIXED(31),
2 L ELB HIGH BIN FIXED(31),
   2 R ELB(1 REFER(L ELB LOW):1 REFER(L ELB HIGH)) BIN FLOAT(24);
DCL 1 S EHM BASED (P EHM),
   2 L_EHM_LOW BIN FIXED(31),
2 L_EHM_HIGH BIN FIXED(31),
   2 R EHM(1 REFER(L EHM LOW):1 REFER(L EHM HIGH))
                                                     BIN FLOAT (24);
DCL 1 S_EHB BASED(P EHB),
   2 L EHB LOW BIN FIXED(31),
   2 L EHB HIGH BIN FIXED(31),
   2 R EHB(1 REFER(L EHB LOW):1 REFER(L EHB HIGH)) BIN FLOAT(24);
DCL 1 S_YLTM BASED(P_YLTM),
   2 L YLTM LOW BIN FIXED(31),
    2 L YLTM HIGH BIN FIXED (31),
   2 R_YLTM(1 REFER(L_YLTM_LOW):1 REFER(L_YLTM_HIGH)) BIN FLOAT(24);
DCL 1 S YLTB BASED (P YLTB),
   2 L YLTB LOW BIN FIXED (31),
   2 L YLTB HIGH BIN FIXED (31),
   2 R YLTB(1 REFER(L YLTB LOW):1 REFER(L YLTB HIGH)) BIN FLOAT(24);
DCL 1 S YLBM BASED(P_YLBM),
   2 L YLBM LOW BIN FIXED (31),
   2 L YLBM HIGH BIN FIXED(31),
    2 R YLBM(1 REFER(L YLBM LOW):1 REFER(L YLBM HIGH)) BIN FLOAT(24);
DCL 1 S YLBB BASED (P YLBB),
   2 L YLBB LOW BIN FIXED(31),
   2 L_YLBB_HIGH BIN FIXED(31),
   2 R YLBB(1 REFER(L YLBB LOW):1 REFER(L YLBB HIGH)) BIN FLOAT(24);
DCL 1 S YHTM BASED (P YHTM),
   2 L YHTM LOW BIN FIXED (31),
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2 L YHTM HIGH BIN FIXED(31),
    2 R YHTM(1 REFER(L YHTM LOW):1 REFER(L YHTM HIGH)) BIN FLOAT(24);
DCL 1 S YHTB BASED (P YHTB),
    2 L YHTB LOW BIN FIXED (31),
    2 L YHTB HIGH BIN FIXED (31),
2 R_YHTB(1 REFER(L_YHTB_LOW):1 REFER(L_YHTB_HIGH))
DCL 1 S_YHBM BASED(P_YHBM),
                                                          BIN FLOAT(24);
    2 L YHBM LOW BIN FIXED (31),
    2 L_YHBM_HIGH BIN FIXED(31),
    2 R YHBM(1 REFER(L YHBM LOW):1 REFER(L YHBM HIGH)) BIN FLOAT(24);
DCL 1 S YHBB BASED (P YHBB),
    2 L_YHBB_LOW BIN FIXED(31),
    2 L YHBB HIGH BIN FIXED(31),
    2 R_YHBB(1 REFER(L_YHBB_LOW):1 REFER(L_YHBB_HIGH)) BIN FLOAT(24);
DCL 1 S PKEV BASED (P PKEV),
    2 L_PKEV_LOW BIN FIXED(31),
2 L_PKEV_HIGH BIN FIXED(31),
    2 R PKEV (1 REFER(L PKEV LOW):1 REFER(L PKEV HIGH)) BIN FLOAT(24);
DCL 1 S PMEV BASED (P PMEV),
    2 L_PMEV_LOW BIN FIXED(31),
2 L_PMEV_HIGH BIN FIXED(31),
    2 R PMEV (1 REFER(L PMEV LOW): 1 REFER(L PMEV HIGH)) BIN FLOAT(24);
DCL 1 S_PLO BASED(P_PLO),
    2 L PLO LOW BIN FIXED(31),
    2 L PLO HIGH BIN FIXED (31),
    2 R PLO(1 REFER(L PLO LOW):1 REFER(L PLO HIGH)) BIN FLOAT(24);
DCL 1 S_PHI BASED(P PHI),
    2 L PHI LOW BIN FIXED (31),
    2 L PHI HIGH BIN FIXED (31),
    2 R PHI(1 REFER(L PHI LOW):1 REFER(L PHI HIGH)) BIN FLOAT(24);
DCL 1 S YELT BASED (P YELT),
    2 L YELT LOW BIN FIXED (31),
    2 L YELT HIGH BIN FIXED (31),
    2 R YELT(1 REFER(L YELT LOW):1 REFER(L YELT HIGH)) BIN FLOAT(24);
DCL 1 S YEHT BASED (P YEHT),
    2 L YEHT LOW BIN FIXED (31),
    2 L YEHT HIGH BIN FIXED (31),
2 R_YEHT(1 REFER(L_YEHT_LOW):1 REFER(L_YEHT_HIGH)) BIN FLOAT(24);
DCL 1 S_YELB BASED(P_YELB),
    2 L YELB LOW BIN FIXED (31),
    2 L_YELB_HIGH BIN FIXED(31),
    2 R YELB(1 REFER(L YELB LOW):1 REFER(L YELB HIGH)) BIN FLOAT(24);
DCL 1 S YEHB BASED (P YEHB),
    2 L_YEHB_LOW BIN FIXED(31),
    2 L YEHB HIGH BIN FIXED (31),
    2 R_YEHB(1 REFER(L_YEHB_LOW):1 REFER(L_YEHB_HIGH)) BIN FLOAT(24);
DCL 1 S EKEV BASED (P EKEV),
    2 L_EKEV_LOW BIN FIXED(31),
2 L_EKEV_HIGH BIN FIXED(31),
    2 R EKEV(1 REFER(L EKEV LOW):1 REFER(L EKEV HIGH)) BIN FLOAT(24);
DCL 1 S EMEV BASED (P EMEV),
    2 L EMEV LOW BIN FIXED (31),
    2 L EMEV HIGH BIN FIXED(31),
    2 R EMEV(1 REFER(L EMEV LOW):1 REFER(L EMEV HIGH)) BIN FLOAT(24);
DCL 1 S_ELO BASED(P_ELO),
    2 L ELO LOW BIN FIXED (31),
    2 L ELO HIGH BIN FIXED(31),
    2 R ELO(1 REFER(L ELO LOW):1 REFER(L ELO HIGH)) BIN FLOAT(24);
DCL 1 S EHI BASED (P EHI),
    2 L EHI LOW BIN FIXED(31),
    2 L EHI HIGH BIN FIXED(31),
    2 R EHI(1 REFER(L EHI LOW):1 REFER(L EHI HIGH)) BIN FLOAT(24);
DCL 1 S E EVR BASED (P E EVR),
    2 L E EVR LOW BIN FIXED (31),
    2 L E EVR HIGH BIN FIXED(31),
    2 R E EVR (1 REFER (L E EVR LOW): 1 REFER (L E EVR HIGH)) BIN FLOAT (24);
DCL 1 S E MOM BASED (P E MOM),
    2 L E MOM LOW BIN FIXED(31),
    2 L_E_MOM_HIGH BIN FIXED(31),
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2 R E MOM(1 REFER(L E MOM LOW):1 REFER(L E MOM HIGH)) BIN FLOAT(24);
DCL 1 S DTEA BASED (P DTEA),
    2 L DTEA LOW BIN FIXED (31),
    2 L DTEA HIGH BIN FIXED (31),
    2 R DTEA(1 REFER(L DTEA LOW):1 REFER(L DTEA HIGH)) BIN FLOAT(24);
DCL 1 S_E_DAU BASED(P_E_DAU),
    2 L E DAU LOW BIN FIXED(31),
    2 L E DAU HIGH BIN FIXED (31),
    2 R_E_DAU(1 REFER(L_E_DAU_LOW):1 REFER(L E DAU HIGH)) BIN FLOAT(24);
DCL 1 S E FISS BASED (P E FISS),
    2 L E FISS LOW BIN FIXED (31),
    2 L E FISS HIGH BIN FIXED(31),
    2 R F FISS (1 REFER(L E FISS LOW): 1 REFER(L E FISS HIGH)) BIN FLOAT(24);
DCL 1 S E EVRF BASED (P E EVRF),
    2 L E EVRF LOW BIN FIXED(31),
    2 L E EVRF HIGH BIN FIXED(31),
                                                                BIN FLOAT(24);
    2 R E EVRF (1 REFER(L E EVRF LOW): 1 REFER(L E EVRF HIGH))
DCL 1 S E PPAC BASED(P E PPAC),
    2 L E PPAC LOW BIN FIXED(31),
    2 L_E_PPAC_HIGH BIN FIXED(31),
    2 R E PPAC (1 REFER(L E PPAC LOW):1 REFER(L E PPAC HIGH)) BIN FLOAT(24);
DCL 1 S NAIL BASED (P NAIL),
    2 L_NAIL_LOW BIN FIXED(31),
    2 L NAIL HIGH BIN FIXED (31),
    2 R NAIL (1 REFER(L NAIL LOW): 1 REFER(L NAIL HIGH)) BIN FLOAT(24);
DCL 1 S NAIH BASED (P NAIH),
    2 L_NAIH_LOW BIN FIXED(31),
2 L NAIH HIGH BIN FIXED(31),
    2 R NAIH (1 REFER(L NAIH LOW): 1 REFER(L NAIH HIGH)) BIN FLOAT(24);
DCL 1 S_RESLSW BASED(P_RESLSW),
    2 L RESLSW LOW
                     BIN FIXED(31),
    2 L RESLSW HIGH BIN FIXED (31),
    2 R RESLSW(1 REFER(L RESLSW LOW):1 REFER(L RESLSW HIGH)) BIN FLOAT(24);
DCL 1 S RESMSW BASED (P RESMSW),
    2 L RESMSW LOW BIN FIXED (31),
    2 L RESMSW HIGH BIN FIXED (31),
    2 R RESMSW(1 REFER(L RESMSW LOW):1 REFER(L RESMSW HIGH))
                                                                BIN FLOAT (24);
DCL 1 S RWSLSW BASED (P RWSLSW),
    2 L RWSLSW LOW BIN FIXED (31),
    2 L RWSLSW HIGH BIN FIXED (31),
    2 R RWSLSW (1 REFER(L RWSLSW LOW):1 REFER(L RWSLSW HIGH))
                                                                BIN FLOAT (24);
DCL 1 S RWSMSW BASED (P RWSMSW),
    2 L RWSMSW LOW BIN FIXED(31),
    2 L RWSMSW HIGH BIN FIXED(31),
    2 R RWSMSW (1 REFER(L RWSMSW LOW): 1 REFER(L RWSMSW HIGH)) BIN FLOAT(24);
DCL 1 S RE BASED (P RE),
    2 L_RE_LOW BIN FIXED(31),
    2 L RE HIGH BIN FIXED (31),
    2 R RE(1 REFER(L RE LOW):1 REFER(L RE HIGH)) BIN FLOAT(24);
DCL 1 S RW BASED (P RW),
    2 L RW LOW BIN FIXED(31),
    2 L RW HIGH BIN FIXED (31),
    2 R RW(1 REFER(L RW LOW):1 REFER(L RW HIGH)) BIN FLOAT(24);
DCL 1 S IPAR BASED (P IPAR),
    2 L IPAR LOW BIN FIXED(31),
2 L IPAR HIGH BIN FIXED(31),
    2 R IPAR(1 REFER(L IPAR LOW):1 REFER(L IPAR HIGH)) BIN FLOAT(24);
DCL 1 S SLOPE BASED (P SLOPE),
    2 L_SLOPE_LOW BIN FIXED(31),
2 L_SLOPE_HIGH BIN FIXED(31),
    2 R SLOPE (1 REFER(L SLOPE LOW):1 REFER(L SLOPE HIGH)) BIN FLOAT(24);
DCL 1 S PPAC BASED (P PPAC),
    2 L PPAC LOW
                   BIN FIXED(31),
    2 L PPAC HIGH BIN FIXED (31),
    2 R PPAC(1 REFER(L PPAC LOW):1 REFER(L PPAC HIGH)) BIN FLOAT(24);
DCL 1 S ERROR BASED (P ERROR),
    2 L ERROR LOW BIN FIXED (31),
    2 L ERROR HIGH BIN FIXED (31),
    2 R_ERROR(1 REFER(L_ERROR_LOW):1 REFER(L_ERROR_HIGH)) BIN FLOAT(24);
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/**** LOCAL PROCEDURE VARIABLES *****/
DCL (P NEXT EVENT, P NEXT SUBEVENT) POINTER;
DCL I
                        BIN FIXED(31);
DCT, J
                      BIN FIXED(31);
DCL K
                      BIN FIXED(31);
DCL L_INCR
                      BIN FIXED(31) INIT(1);
DCL B PAUSE
                        BIT(1) ALIGNED STATIC;
DCL R PRN
                     BIN FLOAT (24) STATIC INIT (1);
                     BIN FLOAT(24) STATIC INIT(1);
DCL R MAX
DCL USLSW
                      BIN FLOAT (53) STATIC;
                     BIN FLOAT(53) STATIC;
DCL USMSW
                      BIN FLOAT (53) STATIC;
DCL USSSW
                     BIN FLOAT(53) STATIC;
DCL MSLSW
DCL MSMSW
                      BIN FLOAT (53) STATIC;
                     BIN FLOAT (53) STATIC;
DCL SLSW
DCL SMSW
                     BIN FLOAT (53) STATIC;
                     BIN FLOAT(53) STATIC;
BIN FLOAT(53) STATIC;
DCL MLSW
DCL MMSW
                    BIN FLOAT(53) STATIC INIT(0.);
DCL TIMEUS
                    BIN FLOAT(53) STATIC INIT(0.);
DCL TIMEMS
DCL TIMESEC
                       BIN FLOAT (53) STATIC INIT (0.);
                     BIN FLOAT(53) STATIC INIT(0.);
DCL TIME
DCL TIMEMIN
                      BIN FLOAT(53) STATIC INIT(0.);
DCL B RE
                      BIT(1) ALIGNED STATIC;
                     BIT(1) ALIGNED STATIC;
DCL B RW
DCL B TIME
                      BIT(1);
                     BIN FLOAT(53) INIT(1000.0000000);
DCL N1000
DCL N216
                     BIN FLOAT(53) INIT(65536.000000);
DCL N224
                      BIN FLOAT (53) INIT (16777216.000);
DCL N232
                     BIN FLOAT (53) INIT (4294967296.0);
                     CHAR (128) VARYING;
BIN FLOAT (24);
DCL C TIME
DCL R EPUNCH
DCL R SPUNCH
                      BIN FLOAT (24);
DCL B MOTHER
                     BIT(1) ALIGNED STATIC;
DCL B_DAUGHTER
DCL B_EVAP
                      BIT(1) ALIGNED STATIC;
                     BIT(1) ALIGNED STATIC;
                      BIT(1) ALIGNED STATIC;
DCL B EVAPF
                       BIT(1) ALIGNED STATIC;
DCL B_FISSION
                     BIT(1) ALIGNED STATIC;
DCL B PPAC
DCL B EVR
                     BIT(1) ALIGNED STATIC;
DCL B MOM
                     BIT(1) ALIGNED STATIC;
DCL B DAU
                       BIT(1) ALIGNED STATIC;
DCL B FISS
                       BIT(1) ALIGNED STATIC;
DCL B EVRF
                      BIT(1) ALIGNED STATIC;
                       BIT(1) ALIGNED STATIC;
DCL B PUNCH(8)
DCL B PUNCHTHROUGH
                       BIT(1) ALIGNED STATIC;
DCL C
                       BIN FIXED(31);
DCL N
                       BIN FIXED(31);
DCL Q
                        BIN FIXED(31) INIT(0);
                       BIN FIXED(31) INIT(0);
DCT<sub>1</sub> R
DCL U
                      BIN FIXED(31) INIT(0);
DCL I_EVCT
                     BIN FIXED(31) STATIC INIT(0);
DCL T EA
                      BIN FLOAT (24) STATIC;
DCL T EA MAX
                     BIN FLOAT(24) STATIC;
                     BIN FLOAT(24) STATIC;
DCL T EA MIN
DCL T_AA
DCL T_AA_MAX
                      BIN FLOAT (24) STATIC;
                     BIN FLOAT(24) STATIC;
DCL T AA MIN
                      BIN FLOAT(24) STATIC;
DCL T_EF
                       BIN FLOAT (24) STATIC;
DCL T EF MAX
                       BIN FLOAT(24) STATIC;
DCL T EF MIN
                      BIN FLOAT(24) STATIC;
DCL C EVAP
                      CHAR (128) VARYING;
DCL P_DIFF_EA
DCL P_DIFF_AA
                      BIN FLOAT (24) STATIC;
                     BIN FLOAT(24) STATIC;
DCL P DIFF EF
                      BIN FLOAT(24) STATIC;
DCL E_RATIO
                       BIN FLOAT (24) STATIC;
```

```
DCL LOGT_EA BIN FLOAT(24) STATIC;
DCL R DT
                            BIN FLOAT (24);
______
      INTERACTIVE PARAMETERS USED IN THIS CODE
       ______
                 DESCRIPTION
PARAMETER
                                                       VALUES USED
R_IPAR(1) DO CORRELATIONS? 1=YES,0=NO
R_IPAR(2) PRINT CORRELATIONS 1=YES,0=NO
R_IPAR(3) NUMBER OF CORRELATIONS VARIABLE
R_IPAR(4) CLEAR BUFFERS/EVENT COUNTERS 1=YES,0=NO
R_IPAR(5) POSITION GATE ALPHAS LOW VARIABLE
R_IPAR(6) POSITION GATE ALPHAS HIGH VARIABLE
R_IPAR(7) POSITION GATE FISSIONS LOW VARIABLE
R_IPAR(8) POSITION GATE FISSIONS HIGH VARIABLE
R_IPAR(9)
                                                       -----
                   _____
                                                    VARIABLE
VARTAPT
R IPAR(9)
R IPAR(10) TIME EVAP-MAX
R IPAR(11) TIME EVAP-MIN
R IPAR(12) TIME ALPHA-MAX
R IPAR(13) TIME ALPHA-MIN
                TIME EVAP-MIN VARIABLE
TIME ALPHA-MAX VARIABLE
TIME ALPHA-MIN VARIABLE
TIME EVAP-FISS MAX VARIABLE
TIME EVAP-FISS MIN VARIABLE
                                                         VARIABLE
R IPAR(14)
R_IPAR(15)
R IPAR(16)
R_IPAR(17)
LIST EVENTS QUESTION
R_IPAR(18)
STRIP LISTER
R_IPAR(19)
MINIMUM EVENT NUMBER
R_IPAR(20)
MAXIMUM NUMBER OF EVENTS
                                                         VARIABLE
                                                           1-32
                                                           VARIABLE
                                                          VARTABLE
R IPAR(21)
TIMES? 1=YES,0=NO
N_IPAR(24) PRINT TIME LOW RANGE VARIABLE
R_IPAR(25) PRINT TIME HIGH RANGE VARIABLE
R_IPAR(26) PRINT EVENTS VICTOR? 1=YES,0=
R_IPAR(27)
R_IPAR(28)
R_IPAR(22)
                                                          1=YES, 0=NO
R_IPAR(29)
               CORRELATED EVENT COUNTER 1=YES, 0=NO
R IPAR(30)
 ______
   X$ANAL - MAIN BODY OF PROCEEDURE STARTS HERE
 ______
 */
 @DCL MSG(XIO NOOUTPUT);
 @ON ANY W(U CLEANUP);
STS$VALUE=1;
P SA$BUFHE = P BUFFER;
                                                       /* SET POINTER TO BUFFER HEADER */
                                                        /* SET POINTER TO EVENT HEADER */
P SA$VE10 1 = P EVENT;
 P NEXT_EVENT = ADDR(LA$VE10_1_NEXT);
 P NEXT SUBEVENT = ADDR(IA$VE10 1(1));
R ERROR(1) = 0.;
 DO WHILE (P NEXT SUBEVENT ^= P NEXT EVENT); /* LOOP OVER SUBEVENTS IN EVENT */
    P_SA$VES10_1 = P_NEXT_SUBEVENT;
                                                      /* SET POINTER TO SUBEVENT HDR */
    P NEXT SUBEVENT = ADDR (LA$VES10 1 NEXT);
```

```
I EVCT=I EVCT+1;
   R ERROR(1) = 0.5;
   R = RROR(2) = 0.5;
                       CALL POSITION; /** POSITION PROC **/
CALL SINGIFE: /**
   R = RROR(1) = 1.;
   R ERROR(1) = 2.;
   R = RROR(1) = 3.;
   R = RROR(1) = 4.;
                                               /** POSITION DETERMINATION **/
                                             /** SINGLES SPECTRA **/
   R = ERROR(1) = 5.;
                                              /** CORRELATION ROUTINE **/
   R_ERROR(1) = 6.; CALL BUFFER;
   R = RROR(1) = 7.;
                         CALL EVENTLIST;
   R = -0.5;
   R = RROR(2) = -0.5;
END;
______
     PROCEDURE FOR UNPACKING RAW DATA INTO RA$SECAM AND MAKING RAW SPECTRA
     SPECTRA-(S)
______
UNPACK RAW : PROCEDURE;
DCL SE INDEX
                         BIN FIXED(31);
/** INITIALIZE RA$SECAM **/ R ERROR(2) = 1.;
DO I = 1 TO 353;
  RA\$SECAM(I)=0.;
R ERROR(2) = 11.;
R = RROR(2) = 12.;
R = RROR(2) = 13.;
R = NAIL(1) = 0.;
R = NAIH(1) = 0.;
R RESLSW(1) = 0.;
R = RESMSW(1) = 0.;
R \text{ RWSLSW}(1) = 0.;
R RWSMSW(1) = 0.;
R RE(1) = 0.;
R = 0.;
\overline{DO} I = 1 TO 32;
   R EL(I) = 0.;
   R \to H(I) = 0.;
   R YLT(I) = 0.;
   R YHT(I) = 0.;
   R YLB(I) = 0.;
   R YHB(I) = 0.;
END;
DO I = 1 TO 16;
   R BL(I) = 0.;
   R BH(I) = 0.;
END;
DO I = 1 TO 8;
   R PT(I) = 0.;
END;
R PPAC(1) = 0.;
R ERROR(2) = 2.;
\overline{\text{IF}} (I EVCT >= R IPAR(19)) & (I EVCT <= R IPAR(20)) & (R IPAR(26) = 1) THEN DO;
   @CALL U$PRTCL(
   'Buf:'|| TRIM(CHAR(LA$BUFHE buf))||
   ', evts:'|| TRIM(CHAR(LA$BUFHE_evt))||
   ', EV len:'||TRIM(CHAR(LA$ve10_1_dlen))||
', t/s:'|| TRIM(CHAR(IA$ve10_1_type))||
   '/'||
                TRIM(CHAR(IA$ve10 1 subtype))||
   ', #:'|| TRIM(CHAR(LA$ve10_1_count))||
'SE len:'|| TRIM(CHAR(LA$ves10_1_dlen))||
   ', t/s:'|| TRIM(CHAR(IA$ves10 1 type))||
   '/'||
                 TRIM(CHAR(IA$ves10_1_subtype))||
```

```
', id:'|| TRIM(CHAR(IA$ves10 1 procid))
   ,U$M prtterm);
END;
/** LOAD RA$SECAM **/ R_ERROR(2) = 3.;
DO I = 1 TO (LA$VES10 1 DLEN - 2) BY 2;
   SE INDEX = FIXED(\overline{IA}$VES10 1(I),31);
   IF (SE INDEX > 0 & SE INDEX <= 354) THEN DO;
      RA$SECAM(SE_INDEX) = POSINT(IA$VES10_1(I+1),1,16);
      IF (R IPAR(\overline{2}6) = 1) & (I EVCT >= R IPAR(19)) & (I EVCT <= R IPAR(20)) THEN DO;
        @CALL U$PRTCL(
        '# '||TRIM(CHAR(IA$ves10 1(I)))||
        ' value '||TRIM(CHAR(IA$ves10 1(I+1)))
        ,U$M prtterm);
      END;
   END:
END;
/** CREATE RAW SPECTRA **/ R ERROR(2) = 4.;
DO I = 1 TO 352;
  IF RA$SECAM(I) > 0 & RA$SECAM(I) < 4095 THEN
$ACCU1(L,DB,$SPECTRUM,S,I,L_INCR,1,RA$SECAM(I));
/** CREATE DATA ELEMENT VALUES **/
R ERROR(2) = 5.;
R ERROR(2) = 6.;
/* EL, EH, YLT, YHT, YLB, YHB */ R ERROR(2) = 7.;
DO I = 1 TO 32;
  IF RA$SECAM(I+31) > 0 & RA$SECAM(I+31) <= 4096 THEN R EL(I) = RA$SECAM(I+31);
   IF RA\$SECAM(I+63) > 0 & RA\$SECAM(I+63) \le 4096 THEN R EH(I) = RA\$SECAM(I+63);
   IF RA\$SECAM(I+95) > 0 & RA\$SECAM(I+95) \le 4096 THEN R\_YLT(I) = RA\$SECAM(I+95);
   IF RA\$SECAM(I+127) > 0 \& RA\$SECAM(I+127) \le 4096 \text{ THEN } R \text{ YHT}(I) = RA\$SECAM(I+127);
   IF RA\$SECAM(I+159) > 0 \& RA\$SECAM(I+159) <= 4096 THEN R_YLB(I) = RA\$SECAM(I+159);
  IF RA$SECAM(I+191) > 0 & RA$SECAM(I+191) <= 4096 THEN R YHB(I) = RA$SECAM(I+191);
/* BACKWARDS & PUNCHTHROUGHS */ R ERROR(2) = 8.;
DO T = 1 TO 16:
   IF RA\$SECAM(I+223) > 0 \& RA\$SECAM(I+223) <= 4096 THEN R BL(I) = RA\$SECAM(I+223);
   IF RA\$SECAM(I+239) > 0 & RA\$SECAM(I+239) \le 4096 THEN RBH(I) = RA\$SECAM(I+239);
END:
DO I = 1 TO 8;
  IF RA\$SECAM(I+271) > 0 & RA\$SECAM(I+271) <= 4096 THEN R PT(I) = RA\$SECAM(I+271);
/* PPAC */ R ERROR(2) = 9.;
IF RA$SECAM(260) > 0 & RA$SECAM(260) <= 4096 THEN R PPAC(1) = RA$SECAM(260);
/* SODIUM IODIDE */
IF RA$SECAM(268) > 0 & RA$SECAM(268) <= 4096 THEN R NAIL(1) = RA$SECAM(268);
IF RA$SECAM(269) > 0 & RA$SECAM(269) <= 4096 THEN R NAIH(1) = RA$SECAM(269);
/* RUTHERFORDS */
IF RA$SECAM(25) > 0 & RA$SECAM(25) <= 4096 THEN R RESLSW(1) = RA$SECAM(25);
IF RA\$SECAM(26) > 0 \& RA\$SECAM(26) \le 4096 THEN R RESMSW(1) = RA\$SECAM(26);
IF RA\$SECAM(27) > 0 \& RA\$SECAM(27) \le 4096 THEN RWSLSW(1) = RA\$SECAM(27);
IF RA$SECAM(28) > 0 & RA$SECAM(28) <= 4096 THEN R_RWSMSW(1) = RA$SECAM(28);
IF RA$SECAM(270) > 0 & RA$SECAM(270) <= 4096 THEN R RE(1) = RA$SECAM(270);
IF RA\$SECAM(271) > 0 \& RA\$SECAM(271) \le 4096 \text{ THEN R } RW(1) = RA\$SECAM(271);
END UNPACK RAW;
______
    TIME ANALYSIS
     SPECTRA-(SEC, TIME, RE RATE, RW RATE, RUTH RATE, CHOPMS, CHOPUS)
```

```
TIMES : PROCEDURE;
/** ZERO PARAMETERS **/
USLSW = 0.;
USMSW = 0.;
USSSW = 0.;
MSLSW = 0.;
MSMSW = 0.;
SLSW = 0.;
SMSW = 0.;
MLSW = 0.;
MMSW = 0.;
TIMEUS = 0.;
TIMEMS = 0.;
TIMESEC = 0.;
TIMEMIN = 0.;
TIME = 0.;
B_RE = '0'B;
B RW = '0'B;
/** DEFINE TIME VARIABLES **/
USLSW = RA$SECAM(3);
USMSW = RA$SECAM(4);
MSLSW = RA$SECAM(5);
MSMSW = RA$SECAM(6);
SLSW = RA$SECAM(7);
SMSW = RA$SECAM(8);
MLSW = RA$SECAM(9);
MMSW = RA$SECAM(10);
TIMESEC = SLSW;
TIMEMS = MSLSW + MSMSW*N216;
B_TIME = '1'B;
I = -1;
DO WHILE (B TIME);
   I = I + \overline{1};
   IF (TIMEMS > (I*N232)/N1000) & (TIMEMS < ((I+1)*N232)/N1000) THEN DO;
      USSSW = I;
      TIMEUS = USLSW + USMSW*N216 + USSSW*N232;
      B_{TIME} = '0'B;
   IF I > 50 THEN B TIME = '0'B;
END;
R ERROR(2) = 3.;
\overline{\text{TIME}} = \overline{\text{TIMEMS}}/(N1000);
TIMEMIN = TIMEMS/(60*N1000);
/** PRINT OUT TIMES **/ R ERROR(2) = 4.;
IF (R IPAR(23) = 1) THEN \overline{DO};
   IF TIMESEC > R_IPAR(24) & TIMESEC < R_IPAR(25) THEN DO;
       PUT STRING(C_TIME) EDIT(
       'Time',TIME,
       ' Min', TIMEMIN,
       ' Sec', TIMESEC,
       ' Millisec', TIMEMS,
       ' Microsec', TIMEUS)
       (A, F(13, 6), A, F(13, 6), A, F(6, 0), A, F(11, 0), A, F(14, 0));
       @CALL U$PRTCL(C_TIME, U$M_PRTTERM);
   END;
END;
$ACCU(L, DB, $SPECTRUM, SEC, L INCR, 1, TIMESEC);
$ACCU(L,DB,$SPECTRUM,TIME,L_INCR,1,TIME);
$ACCU(L, DB, $SPECTRUM, TIMEMIN, L_INCR, 1, TIMEMIN);
/** RUTHERFORD RATE SPECTRA **/ R ERROR(2) = 5.;
```

```
$COND(WC,DB,$CONDITION,RE,B RE,1,R RE(1));
$COND(WC,DB,$CONDITION,RW,B RW,1,R RW(1));
IF B RE THEN DO;
   $\overline{A}CCU(L,DB,$SPECTRUM,RE RATE,L INCR,1,TIME);
   $ACCU(L,DB,$SPECTRUM,RUTH RATE,L INCR,1,TIME);
IF B RW THEN DO;
   $ACCU(L, DB, $SPECTRUM, RW RATE, L INCR, 1, TIME);
   $ACCU(L,DB,$SPECTRUM,RUTH_RATE,L_INCR,1,TIME);
END TIMES;
     CALIBRATION ROUTINE AND SP, SUM, SUMP SPECTRA
     SPECTRA-(TMP, SP, SPSUM, PPAC, EVR LOW)
______
*/
CALIBRATION : PROCEDURE;
/** ZERO PARAMETERS **/
B PAUSE = '0'B;
B PPAC = '0'B;
R PRN = 0.;
\overline{DO} I = 1 TO 32;
   R EKEV(I) = 0.;
   R = EMEV(I) = 0.;
END;
R ELO(1) = 0.;
R_EHI(1) = 0.;
/** TMP CREATION LINE AND PAUSE CONDITION **/
IF RA\$SECAM(1) > 0 \& RA\$SECAM(1) < 30001 THEN DO;
   $ACCU(L,DB,$SPECTRUM,TMP,L INCR,1,RA$SECAM(1));
   $COND(WC,DB,$CONDITION,PAUSE,B PAUSE,1,RA$SECAM(1));
/** SP SPECTRA CREATION **/
DO I = 1 TO 32;
   IF B PAUSE THEN DO;
      \overline{IF} R EL(I) > 150 THEN DO;
         $ACCU1(L,DB,$SPECTRUM,SP,I,L_INCR,1,R_EL(I));
         $ACCU(L, DB, $SPECTRUM, SPSUM, L INCR, 1, R EL(I));
      END:
   END;
END;
/** GENERATE RANDOM NUMBER **/
R PRN=U$RANDOM(R MAX);
/** FILL IN ARRAY WITH CALIBRATED DATA **/
DO I = 1 TO 32;
   IF R EL(I) > 0 THEN DO;
      R 	ext{ EKEV(I) = ((R 	ext{ EL(I) + R PRN) *R ELM(I)) + R ELB(I);}}
      R = KEV(I) = R = KEV(I) * (1.0 - R SLOPE(I) * ((R YLT(I)/R EL(I)) - 0.5));
      \overline{IF} (R EKEV(\overline{I}) > 250) & (R_EKEV(\overline{I}) > R_ELO(\overline{I}) THEN R_ELO(\overline{I}) = R_EKEV(\overline{I});
END;
DO I = 1 TO 32;
   IF R EH(I) > 0 THEN DO;
      R = EMEV(I) = R EH(I);
      \overline{\text{IF}} (R EMEV(\overline{\text{I}}) > 50) & (R EMEV(I) > R EHI(1)) THEN R EHI(1) = R EMEV(I);
   END;
END;
/** PPAC CONDITION SPECTRA **/
$COND(WC,DB,$CONDITION,PPAC,B_PPAC,1,R_PPAC(1));
```

```
IF B PPAC THEN DO;
   $\overline{ACCU(L,DB,$SPECTRUM,PPAC,L INCR,1,R PPAC(1));}
   IF R_ELO(1) > 5 THEN DO;
      $ACCU(L,DB,$SPECTRUM,EVR LOW,L INCR,1,R ELO(1));
END:
END CALIBRATION;
______
     POSITION DETERMINATION
     SPECTRA-(POS T, POS B)
POSITION : PROCEDURE;
/** ZERO PARAMETERS **/
R SEL(1) = 0.;
R \subset EL(1) = 0.;
R_SEH(1) = 0.;
R = 0.;
R SYLT(1) = 0.;
R CYLT(1) = 0.;
R SYLB(1) = 0.;
R CYLB(1) = 0.;
R = SYHT(1) = 0.;
R CYHT(1) = 0.;
R SYHB(1) = 0.;
R CYHB(1) = 0.;
R SBL(1) = 0.;
R \subset BL(1) = 0.;
R = 0.;
R \subset BH(1) = 0.;
R SPT(1) = 0.;
R CPT(1) = 0.;
\overline{DO} I = 1 TO 32;
   R PKEV(I) = 0.;
   R_PMEV(I) = 0.;
   R YELT(I) = 0.;
   R YEHT(I) = 0.;
   R YELB(I) = 0.;
   R YEHB(I) = 0.;
END;
R PLO(1) = 0.;
R PHI(1) = 0.;
/** STRIP AND MAX CHANNEL ROUTINE **/
DO I = 1 TO 32;
   IF (R EL(I) > 50) & (R EL(I) > R CEL(1)) THEN R SEL(1) = I;
   IF (R EL(I) > 50) \& (R EL(I) > R CEL(1)) THEN R CEL(1) = R EL(I);
   IF (R EH(I) > 50) & (R EH(I) > R CEH(1)) THEN R SEH(1) = I;
    \label{eq:ceh}  \text{IF } (\text{R\_EH}(\text{I}) \ > \ 50) \ \& \ (\text{R\_EH}(\text{I}) \ > \ \text{R\_CEH}(\text{I})) \ \ \text{THEN } \ \text{R\_CEH}(\text{I}) \ = \ \text{R\_EH}(\text{I}); 
   IF (R_YLT(I) > 50) & (R_YLT(I) > R_CYLT(1)) THEN R_SYLT(1) = I;
   IF (R YLT(I) > 50) & (R YLT(I) > R CYLT(1)) THEN R CYLT(1) = R YLT(I);
   IF (R_YLB(I) > 50) & (R_YLB(I) > R_CYLB(1)) THEN R_SYLB(1) = I;
   IF (R\_YLB(I) > 50) & (R\_YLB(I) > R\_CYLB(1)) THEN R\_CYLB(1) = R\_YLB(I); IF (R\_YHT(I) > 50) & (R\_YHT(I) > R\_CYHT(1)) THEN R\_SYHT(1) = I;
   IF (R YHT(I) > 50) \& (R YHT(I) > R CYHT(1)) THEN R CYHT(1) = R YHT(I);
   IF (R_YHB(I) > 50) & (R_YHB(I) > R_CYHB(1)) THEN R_SYHB(1) = I;
   IF (R YHB(I) > 50) \& (R YHB(I) > R CYHB(1)) THEN R CYHB(1) = R YHB(I);
END;
DO I = 1 TO 16;
   IF (R BL(I) > 50) & (R BL(I) > R CBL(1)) THEN R SBL(1) = I;
   IF (R BL(I) > 50) \& (R BL(I) > R CBL(1)) THEN R CBL(1) = R BL(I);
   IF (R_BH(I) > 50) \& (R_BH(I) > R_CBH(1)) THEN R_SBH(1) = I;
```

```
IF (R BH(I) > 50) \& (R BH(I) > R CBH(1)) THEN R CBH(1) = R BH(I);
END:
DO I = 1 TO 8;
  IF (R PT(I) > 50) \& (R PT(I) > R CPT(1)) THEN R SPT(1) = I;
  IF (R PT(I) > 50) & (R PT(I) > R CPT(1)) THEN R CPT(1) = R PT(I);
/** POSITION LOW **/
IF R CEL(1) > 0 THEN DO;
  R \text{ PLO}(1) = (R \text{ CYLT}(1) - R \text{ CYLB}(1)) / (R \text{ CEL}(1)) * 500;
   $ACCU(L,DB,$SPECTRUM,PLO,L_INCR,1,R_PLO(1));
END;
/** POSITION HIGH **/
IF R CEH(1) > 0 THEN DO;
   R = PHI(1) = (R CYHT(1) - R CYHB(1)) / (R CEH(1)) * 500;
   $ACCU(L,DB,$SPECTRUM,PHI,L_INCR,1,R_PHI(1));
END:
END POSITION;
    SINGLES SPECTRA CREATION
    SPECTRA-(E KEV, SUM, E MEV, SUMMEV, EP KEV, SUMP, EA KEV, SUMA
            RUTHEAST, RUTHWEST, EVR HIGH)
______
SINGLES : PROCEDURE;
/** ZERO PARAMETERS **/
DO I = 1 TO 32;
  IF R EL(I) > 50 \& R EL(I) < 4095 THEN
$ACCU1(L,DB,$SPECTRUM,EKEV,I,L INCR,1,R EKEV(I));
IF R ELO(1) > 5 THEN $ACCU(L,DB,$SPECTRUM,SUML,L INCR,1,R ELO(1));
DO I = 1 TO 32;
  IF R EH(I) > 50 & R EH(I) < 4095 THEN
$ACCU1(L,DB,$SPECTRUM,EMEV,I,L INCR,1,R EMEV(I));
IF R EHI(1) > 5 THEN $ACCU(L,DB,$SPECTRUM,SUMH,L INCR,1,R EHI(1));
IF B PAUSE THEN DO;
  DO I = 1 TO 32;
     IF R EL(I) > 50 & R EL(I) < 4095 THEN
$ACCU1(L,DB,$SPECTRUM,EPKEV,I,L INCR,1,R EKEV(I));
  END:
   IF R ELO(1) > 5 THEN $ACCU(L,DB,$SPECTRUM,SUMLP,L INCR,1,R ELO(1));
   DO I = 1 TO 32;
     IF R EH(I) > 50 \& R EH(I) < 4095 THEN
$ACCU1(L,DB,$SPECTRUM,EPMEV,I,L_INCR,1,R_EMEV(I));
  END;
   IF R EHI(1) > 5 THEN $ACCU(L,DB,$SPECTRUM,SUMHP,L INCR,1,R EHI(1));
END;
IF ^B PPAC THEN DO;
   DO^{-}I = 1 TO 32;
     IF R EL(I) > 50 & R EL(I) < 4095 THEN
$ACCU1(L,DB,$SPECTRUM,EAKEV,I,L INCR,1,R EKEV(I));
   IF R ELO(1) > 5 THEN $ACCU(L,DB,$SPECTRUM,SUMLA,L INCR,1,R ELO(1));
  DO I = 1 TO 32;
```

```
IF R EH(I) > 50 & R EH(I) < 4095 THEN
ACCU1(L,DB,SPECTRUM,EAMEV,I,L_INCR,1,R_EMEV(I));
   IF R EHI(1) > 5 THEN $ACCU(L,DB,$SPECTRUM,SUMHA,L INCR,1,R EHI(1));
END;
$ACCU(L,DB,$SPECTRUM,RUTHEAST,L INCR,1,R RE(1));
$ACCU(L, DB, $SPECTRUM, RUTHWEST, L INCR, 1, R RW(1));
END SINGLES;
     BUFFER ROUTINE
     SPECTRA-(DT EA, DP EA, E EVAP, E MOTHER, P EVAP, P MOTHER
             DT_AA, DP_AA, E_EVAPALPHA, E_DAUGHTER, P_EVAPALPHA, P_DAUGHTER, E_RATIO
              DT EF, DP EF, E EVAPF, E FISSION, P EVAPF, P FISSION)
______
*/
BUFFER : PROCEDURE;
/** ZERO PARAMETERS **/
T EA MAX = 0.;
T = A MIN = 0.;
T AA MAX = 0.;
T_AA_MIN = 0.;
T EF MAX = 0.;
T = T = MIN = 0.;
B MOTHER = '0'B;
B DAUGHTER = '0'B;
B_FISSION = '0'B;
B EVAP = '0'B;
B EVAPF = '0'B;
B MOM = '0'B;
B DAU = '0'B;
B EVR = '0'B;
B = VRF = '0'B;
B FISS = '0'B;
\overline{DO} I = 1 TO 8;
  B PUNCH(I) = '0'B;
END;
B PUNCHTHROUGH = '0'B;
T EA = 0.;
T_AA = 0.;
T_EF = 0.;
LOGT EA = 0.;
P DIFF EA = 0.;
P DIFF_AA = 0.;
P DIFF EF = 0.;
R E EVR(1) = 0.;
R = MOM(1) = 0.;
R DTEA(1) = 0.;
R = DAU(1) = 0.;
R_E^-FISS(1) = 0.;
R = EVRF(1) = 0.;
R EPUNCH = 0.;
R SPUNCH = 0.;
/** PPAC ISSUE **/
IF R_PPAC(1) < 100 & R_PPAC(1) > 0 THEN R_PPAC(1) = 0;
/** SET CONDITIONS **/
```

```
$COND(WC,DB,$CONDITION,E EVR,B EVR,1,R CEL(1));
$COND(WC,DB,$CONDITION,E_MOM,B_MOM,1,R_CEL(1));
$COND(WC, DB, $CONDITION, E_DAU, B_DAU, 1, R_CEL(1));
$COND(WC, DB, $CONDITION, E_FISS, B_FISS, 1, R_CEH(1));
$COND(WC,DB,$CONDITION,E EVRF,B EVRF,1,R CEL(1));
DO I = 1 TO 8;
   IF R PT(I) > R EPUNCH THEN DO;
      R = PUNCH = R PT(I);
     RSPUNCH = I;
   END;
   $COND1(WC,DB,$CONDITION,PUNCH,I,B PUNCH(I),1,R_PT(I));
END;
IF B PUNCH(1) | B PUNCH(2) | B PUNCH(3) | B PUNCH(4) | B PUNCH(5) |
  B PUNCH(6) | B PUNCH(7) | B PUNCH(8) THEN B PUNCHTHROUGH = '1'B;
IF 'B PUNCHTHROUGH THEN DO;
   $ACCU(L,DB,$SPECTRUM,SUMAPT,L INCR,1,R ELO(1));
END;
IF B PUNCHTHROUGH THEN DO;
   $ACCU(L,DB,$SPECTRUM,SUMPT,L INCR,1,R ELO(1));
END;
/** SET TIMING MAXIMUMS AND MINIMUMS **/
T_EA_MAX = R_IPAR(10);
T = A MIN = R IPAR(11);
TAAMAX = RIPAR(12);
T AA MIN = R IPAR(13);
T_{EF}MAX = R_{IPAR(14)};
T EF MIN = R IPAR(15);
/** DECLARE BUFFERS **/
DCL 1 EDATA (4000) STATIC,
                       BIN FLOAT (53),
    2 Т
    2 E
                       BIN FLOAT (24),
    2 S
                       BIN FLOAT (24),
    2 P
                        BIN FLOAT (24),
    2 EV
                       BIN FLOAT(24),
    2 TMP
                       BIN FLOAT (24),
    2 PPAC
                        BIN FLOAT (24);
DCL (ECOUNTER, ELAST) BIN FIXED(31) STATIC INIT(0);
DCL 1 MDATA(4000) STATIC,
    2 T
                        BIN FLOAT (53),
    2 E
                       BIN FLOAT (24),
    2 S
                        BIN FLOAT (24),
    2 P
                         BIN FLOAT (24),
    2 EV
                        BIN FLOAT (24),
    2 TMP
                       BIN FLOAT(24),
    2 PPAC
                       BIN FLOAT(24);
DCL (MCOUNTER, MLAST) BIN FIXED(31) STATIC INIT(0);
DCL 1 DDATA(4000) STATIC,
    2 T
                        BIN FLOAT (53),
    2 E
                       BIN FLOAT (24),
    2 S
                       BIN FLOAT (24),
    2 P
                        BIN FLOAT(24),
    2 EV
                      BIN FLOAT (24),
    2 TMP
                      BIN FLOAT (24),
    2 PPAC
                       BIN FLOAT(24);
DCL (DCOUNTER, DLAST) BIN FIXED (31) STATIC INIT(0);
DCL 1 EFDATA(4000) STATIC,
                        BIN FLOAT (53),
   2 T
    2 E
                       BIN FLOAT(24),
                       BIN FLOAT(24),
    2 S
    2 P
                        BIN FLOAT (24),
    2 EV
                       BIN FLOAT (24),
    2 TMP
                       BIN FLOAT (24),
    2 PPAC
                        BIN FLOAT (24);
DCL (EFCOUNTER, EFLAST) BIN FIXED(31) STATIC INIT(0);
```

```
DCL 1 FDATA(4000) STATIC,
    2 T
                        BIN FLOAT (53),
    2 E
                       BIN FLOAT(24),
    2 S
                        BIN FLOAT (24),
    2 P
                         BIN FLOAT (24),
    2 EV
                        BIN FLOAT(24),
    2 TMP
                       BIN FLOAT (24),
    2 PPAC
                       BIN FLOAT(24);
DCL (FCOUNTER, FLAST) BIN FIXED (31) STATIC INIT(0);
/** CLEAR THE EVENT COUNTERS AND CLEAR THE BUFFERS **/
IF R IPAR(4) = 1 THEN DO;
  ECOUNTER = 0.;
  ELAST = 0.;
  MCOUNTER = 0.;
  MLAST = 0.;
  DCOUNTER = 0.;
   DLAST = 0.;
  EFCOUNTER = 0.;
  EFLAST = 0.;
  FCOUNTER = 0.;
  FLAST = 0.;
  Q = 0.;
  \tilde{R} = 0.;
  U = 0.;
  C = 0.;
  N = 0.;
   I EVCT = 0.;
   \overline{DO} I = 1 TO 4000;
      EDATA(I).T=0.;
      EDATA(I).E=0.;
      EDATA(I).S=0.;
      EDATA(I).P=0.;
      EDATA(I).EV=0.;
      EDATA(I).TMP=0.;
      EDATA(I).PPAC=0.;
      MDATA(I).T=0.;
     MDATA(I).E=0.;
     MDATA(I).S=0.;
     MDATA(I).P=0.;
     MDATA(I).EV=0.;
     MDATA(I).TMP=0.;
     MDATA(I).PPAC=0.;
      DDATA(I).T=0.;
      DDATA(I).E=0.;
      DDATA(I).S=0.;
      DDATA(I).P=0.;
      DDATA(I).EV=0.;
      DDATA(I).TMP=0.;
      DDATA(I).PPAC=0.;
      EFDATA(I).T=0.;
      EFDATA(I).E=0.;
      EFDATA(I).S=0.;
      EFDATA(I).P=0.;
      EFDATA(I).EV=0.;
      EFDATA(I).TMP=0.;
      EFDATA(I).PPAC=0.;
      FDATA(I).T=0.;
      FDATA(I).E=0.;
      FDATA(I).S=0.;
      FDATA(I).P=0.;
      FDATA(I).EV=0.;
      FDATA(I).TMP=0.;
      FDATA(I).PPAC=0.;
   END;
   R_IPAR(4) = 0;
```

```
/** CHECK IF EVENT IS ALPHA OR EVR **/
IF B EVR & (R IPAR(1) = 1) & (B PPAC) & (ABS(R PLO(1)) > 0) THEN B EVAP='1'B;
IF B MOM & (R IPAR(1) = 1) & (^B PPAC) & (ABS(R PLO(1)) > 0) THEN B MOTHER='1'B;
IF B DAU & (R IPAR(1) = 1) & (^B PPAC) & (ABS(R PLO(1)) > 0) THEN B DAUGHTER='1'B;
IF B FISS & (R IPAR(1) = 1) & (^B PPAC) & (ABS(R PHI(1)) > 0) THEN B FISSION='1'B;
IF B EVRF & (R IPAR(1) = 1) & (B PPAC) & (ABS(R PLO(1)) > 0) THEN B EVAPF='1'B;
/** STORAGE OF EVR **/
IF B_EVAP THEN DO;
   I = R SEL(1);
   IF (R^{T}PAR(30) = 1) THEN ACCU1(L, DB, SPECTRUM, CT EVR, I, L INCR, 1, R ELO(1));
   ELAST = ELAST + 1;
   IF ELAST > 4000 THEN ELAST = 1;
   EDATA (ELAST) .T = TIMEMS;
   EDATA (ELAST) .E = R ELO(1);
   EDATA (ELAST) .S = R SEL(1);
   EDATA(ELAST).P = R PLO(1);
   EDATA (ELAST) .EV = \overline{I} EVCT;
   EDATA (ELAST) . TMP = RA$SECAM(1);
   EDATA (ELAST) . PPAC = R PPAC(1);
  IF ECOUNTER < 4000 THEN ECOUNTER = ECOUNTER + 1;
END;
/** STORAGE OF EVR FISSION **/
IF B EVAPF THEN DO;
   I = R SEL(1);
   IF (R IPAR(30) = 1) THEN $ACCU1(L,DB,$SPECTRUM,CT EVRF,I,L INCR,1,R ELO(1));
   EFLAST = EFLAST + 1;
   IF EFLAST > 4000 THEN EFLAST = 1;
   EFDATA (EFLAST) .T = TIMEMS;
   EFDATA(EFLAST).E = R ELO(1);
  EFDATA(EFLAST).S = R SEL(1);
   EFDATA(EFLAST).P = R PLO(1);
   EFDATA (EFLAST) .EV = \overline{I} EVCT;
  EFDATA (EFLAST) . TMP = \overline{RA}$SECAM(1);
   EFDATA(EFLAST).PPAC = R PPAC(1);
  IF EFCOUNTER < 4000 THEN EFCOUNTER = EFCOUNTER + 1;
/** CHECK IF ALPHA CORELATED WITH EVAP RESIDUE **/
IF B MOTHER THEN DO;
   I = R SEL(1);
   IF (R IPAR(30) = 1) THEN $ACCU1(L,DB,$SPECTRUM,CT MOM,I,L INCR,1,R ELO(1));
   0=1;
  MLAST = MLAST + 1;
   IF MLAST > 4000 THEN MLAST = 1;
   MDATA (MLAST) .T = TIMEMS;
  MDATA(MLAST).E = R ELO(1);
  MDATA(MLAST).S = R SEL(1);
  MDATA(MLAST).P = R PLO(1);
   MDATA (MLAST) .EV = I EVCT;
   MDATA(MLAST).TMP = RA$SECAM(1);
   MDATA(MLAST).PPAC = R PPAC(1);
   IF MCOUNTER < 4000 THEN MCOUNTER = MCOUNTER + 1;
   C = ECOUNTER;
   N = ELAST;
   LOOKUPREA: DO WHILE (C > 0);
      T EA = TIMEMS - EDATA(N).T;
      IF (T EA >= T EA MAX) THEN LEAVE LOOKUPREA;
      IF R_SEL(1) = 0 THEN LEAVE LOOKUPREA;
      P DIFF EA = EDATA(N).P-R PLO(1);
      IF (T EA > T EA MIN) &
         (T_EA < T_EA_MAX) & R ELO(1) > 200. &
         (EDATA(N).S = MDATA(MLAST).S) &
         (P_DIFF_EA > R_IPAR(5)) & (P_DIFF_EA < R_IPAR(6)) THEN DO;
         LOGT EA = LOG10 (T EA);
         IF R_{IPAR}(2) = 1 THEN DO;
```

```
PUT STRING(C EVAP) EDIT(
                          'EM-Evap ',
                           ' St', EDATA(N).S,
                          ' Pos', EDATA(N).P,
                          ' Time', EDATA(N).T,
                          ' E Evap', EDATA(N).E,
                           ' Ev Evap', EDATA(N).EV,
                          ' Dp', P DIFF EA,
                           ' Dt',T_EA)
                           (A, A, F(\overline{3}), A, F(10), A, F(10), A, F(6), A, F(8), A, F(12, 3), A, F(8));
                          @CALL U$PRTCL(C EVAP, U$M PRTTERM);
                          PUT STRING(C_EVAP) EDIT(
                                Moth ',
                          ' St', R_SEL(1),
                          ' Pos', R_PLO(1),
' Time', TIMEMS,
                          ' E_Moth', R_ELO(1),
' Ev_Moth', I_EVCT)
                           (A,A,F(3),A,F(10),A,F(10),A,F(6),A,F(8));
                          @CALL U$PRTCL(C EVAP, U$M PRTTERM);
                   END;
                    $ACCU(L, DB, $SPECTRUM, DT EA, L INCR, 1, T EA);
                    $ACCU(L,DB,$SPECTRUM,LOGDT_EA,L_INCR,1,LOGT_EA);
                    $ACCU(L,DB,$SPECTRUM,E EVAP,L INCR,1,EDATA(N).E);
                    $ACCU(L,DB,$SPECTRUM,E MOTHER,L INCR,1,R ELO(1));
                    $ACCU(L,DB,$SPECTRUM,P MOTHER,L INCR,1,R PLO(1));
                    $ACCU(L,DB,$SPECTRUM,P_EVAP,L_INCR,1,EDATA(N).P);
                    $ACCU(L,DB,$SPECTRUM,DP EA,L INCR,1,P DIFF EA);
                    $ACCU(L, DB, $SPECTRUM, RATE EA, L INCR, 1, TIMESEC);
                   R E EVR(1) = EDATA(N).E;
                   R = MOM(1) = R ELO(1);
                   R = PPAC(1) = EDATA(N).PPAC;
                   R DTEA(1) = T EA;
                   0=0+1;
                   IF Q > R_IPAR(3) THEN LEAVE LOOKUPREA;
             N = N - 1;
             IF N <= 0 THEN N = 4000;
             C = C - 1:
      END LOOKUPREA;
END:
/** CHECK IF ALPHA CORRELATED TO ALPHA **/
IF B DAUGHTER THEN DO;
      I = R SEL(1);
       \begin{tabular}{ll}  \begin
      R=1;
      DLAST = DLAST + 1;
      IF DLAST > 4000 THEN DLAST = 1;
      DDATA (DLAST) .T = TIMEMS;
      DDATA (DLAST) .E = R ELO(1);
      DDATA (DLAST) .S = R_SEL(1);
      DDATA (DLAST) .P = R PLO(1);
      DDATA (DLAST) .EV = \overline{I} EVCT;
      DDATA (DLAST) . TMP = \overline{RA}$SECAM(1);
      DDATA (DLAST) . PPAC = R PPAC(1);
      IF DCOUNTER < 4000 THEN DCOUNTER = DCOUNTER + 1;
      C = MCOUNTER;
      N = MLAST;
      LOOKUPALAL: DO WHILE (C > 0);
             T AA = TIMEMS - MDATA(N).T;
             IF (T AA >= T AA MAX) THEN LEAVE LOOKUPALAL;
             IF R \overline{SEL}(1) = 0 THEN LEAVE LOOKUPALAL;
             P DIFF AA = MDATA(N).P-R PLO(1);
             IF (T AA > T AA MIN) &
                    (T_AA < T_AA_MAX) &
                   R = ELO(1) > 200. &
                    (MDATA(N).S = DDATA(DLAST).S) &
                    (P_DIFF_AA > R_IPAR(5)) &
```

```
(P DIFF AA < R IPAR(6)) THEN DO;
          IF R IPAR(2) = 1 THEN DO;
             PUT STRING(C EVAP) EDIT(
             'MD-Moth ',
             ' St', MDATA(N).S,
             ' Pos', MDATA(N).P,
' Time', MDATA(N).T,
             ' E Moth', MDATA(N).E,
              ' Ev_Moth', MDATA(N).EV,
              ' Dp', P DIFF AA,
              ' Dt',T AA)
              (A,A,F(3),A,F(10),A,F(10),A,F(6),A,F(8),A,F(12,3),A,F(8));
             @CALL U$PRTCL(C EVAP, U$M PRTTERM);
             PUT STRING(C EVAP) EDIT(
              ' Daug ',
             ' St', R_SEL(1),
             ' Pos', R_PLO(1),
' Time', TIMEMS,
             ' E_Daug', R_ELO(1),
              ' Ev Daug', I_EVCT)
              (A, A, F(3), A, F(10), A, F(10), A, F(6), A, F(8));
             @CALL U$PRTCL(C EVAP, U$M PRTTERM);
          END;
          E RATIO = (R ELO(1) / MDATA(N).E) * 1000.;
          $ACCU(L,DB,$SPECTRUM,E RATIO,L INCR,1,E RATIO);
          $ACCU(L,DB,$SPECTRUM,DT AA,L INCR,1,T AA);
          $ACCU(L,DB,$SPECTRUM,E_EVAPALPHA,L_INCR,1,MDATA(N).E);
          $ACCU(L,DB,$SPECTRUM,E DAUGHTER,L INCR,1,R ELO(1));
          $ACCU(L,DB,$SPECTRUM,DP AA,L INCR,1,P DIFF AA);
          $ACCU(L,DB,$SPECTRUM,P_DAUGHTER,L_INCR,1,R_PLO(1));
          $ACCU(L,DB,$SPECTRUM,P_EVAPALPHA,L_INCR,1,MDATA(N).P);
          R \in MOM(1) = MDATA(N).E;
          R E DAU(1) = R ELO(1);
          R=R+1;
          IF R > R IPAR(3) THEN LEAVE LOOKUPALAL;
      N = N - 1;
      IF N <= 0 THEN N = 4000;
      C = C - 1;
   END LOOKUPALAL;
END:
/** CHECK IF FISSION CORELATED WITH EVAP RESIDUE **/
IF B FISSION THEN DO;
   I = R SEH(1);
    \texttt{IF} \ (\texttt{R}^{-}\texttt{IPAR}(30) \ = \ 1) \ \texttt{THEN} \ \$\texttt{ACCU1}(\texttt{L}, \texttt{DB}, \$\texttt{SPECTRUM}, \texttt{CT}_\texttt{FISS}, \texttt{I}, \texttt{L}_\texttt{INCR}, 1, \texttt{R}_\texttt{EHI}(1)); 
   U=1;
   FLAST = FLAST + 1;
   IF FLAST > 4000 THEN FLAST = 1;
   FDATA(FLAST).T = TIMEMS;
   FDATA(FLAST).E = R EHI(1);
   FDATA(FLAST).S = R\_SEH(1);
   FDATA(FLAST).P = R PHI(1);
   FDATA (FLAST) .EV = \overline{I} EVCT;
   FDATA(FLAST).TMP = RA$SECAM(1);
   FDATA (FLAST). PPAC = R PPAC(1);
   IF FCOUNTER < 4000 THEN FCOUNTER = FCOUNTER + 1;
   C = EFCOUNTER;
   N = EFLAST;
   LOOKUPEF: DO WHILE (C > 0);
      T EF = TIMEMS - EFDATA(N).T;
      IF (T EF >= T EF MAX) THEN LEAVE LOOKUPEF;
      IF R \overline{SEH}(1) = 0 THEN LEAVE LOOKUPEF;
      P DIFF EF = EFDATA(N).P-FDATA(FLAST).P;
      IF (T EF > T EF MIN) &
          (T_EF < T_EF_MAX) &
          (EFDATA(N).S = FDATA(FLAST).S) &
          (P DIFF EF > R IPAR(7)) &
          (P_DIFF_EF < R_IPAR(8)) THEN DO;
```

```
IF R IPAR(2) = 1 THEN DO;
            PUT STRING (C EVAP) EDIT (
            'EF-Evap ',
            ' St', EFDATA(N).S,
            ' Pos', EFDATA(N).P,
            ' Time', EFDATA(N).T,
             ' E Evap', EFDATA(N).E,
            ' Ev Evap', EFDATA(N).EV,
            ' Dp',P_DIFF_EF,
            ' Dt', T EF)
            (A, A, F(\overline{3}), A, F(10, 3), A, F(10), A, F(6), A, F(8), A, F(12, 3), A, F(9, 3));
            @CALL U$PRTCL(C_EVAP, U$M_PRTTERM);
            PUT STRING(C EVAP) EDIT(
            ' Fiss ',
            ' St', R_SEH(1),
             ' Pos', R_PHI(1),
             ' Time', TIMEMS,
             ' E_Fiss', R_EHI(1),
            ' Ev Fiss', I EVCT)
            (A, A, F(3), A, F(10, 3), A, F(10), A, F(6), A, F(8));
            @CALL U$PRTCL(C_EVAP, U$M_PRTTERM);
         $ACCU(L,DB,$SPECTRUM,DT_EF,L_INCR,1,T_EF);
         $ACCU(L,DB,$SPECTRUM,E EVAPF,L INCR,1,EFDATA(N).E);
         $ACCU(L,DB,$SPECTRUM,E_FISSION,L_INCR,1,R_EHI(1));
         $ACCU(L,DB,$SPECTRUM,P FISSION,L INCR,1,R PHI(1));
         $ACCU(L,DB,$SPECTRUM,P_EVAPF,L_INCR,1,EFDATA(N).P);
         $ACCU(L, DB, $SPECTRUM, DP EF, L INCR, 1, P DIFF EF);
         $ACCU(L,DB,$SPECTRUM,RATE EF,L INCR,1,TIMESEC);
         R E EVRF(1) = EFDATA(N).E;
         R = FISS(1) = R EHI(1);
         U=U+1:
         IF U > R IPAR(3) THEN LEAVE LOOKUPEF;
      END;
      N = N - 1;
      IF N \leq 0 THEN N = 4000;
      C = C - 1;
   END LOOKUPEF;
END:
END BUFFER;
______
    EVENT LISTING PROCEDURE
______
EVENTLIST : PROCEDURE;
IF R IPAR(17) > 10 THEN R SEL(1) = 50;
IF (R IPAR(17) = 1) THEN DO;
   \label{eq:continuous} \mbox{IF I\_EVCT} \ > \ \mbox{R\_IPAR}(19) \ \& \ \mbox{I\_EVCT} \ < \ (\mbox{R\_IPAR}(19) + \mbox{R} \ \mbox{IPAR}(20)) \ \mbox{THEN DO;}
      IF (R SEH(1) = R IPAR(18)) | (R SEL(1) = R IPAR(18)) THEN DO;
         PUT STRING(C EVAP) EDIT(
         'Ev', I_EVCT,
         ' StL', R_SEL(1),
' StH', R_SEH(1),
         ' TIMEMS, TIME,
         ' P_Lo', R_PLO(1),
         ' E Lo', R ELO(1),
         ' P Hi', R PHI(1),
         ' E_Hi', R_EHI(1))
         (A, \overline{F}(8), A, \overline{F}(3), A, F(3), A, F(10, 3), A, F(8, 2), A, F(10, 3), A, F(8, 2), A, F(5));
         @CALL U$PRTCL(C EVAP, U$M PRTTERM);
   END:
```

```
END;
IF (R IPAR(17) = 2) THEN DO;
   IF I EVCT < R IPAR(19) & I EVCT > (R IPAR(19)-R IPAR(20)) THEN DO;
       PUT STRING (C EVAP) EDIT (
       'Ev',I_EVCT,
       ' St', R SEL(1),
       ' Time', TIME,
       ' PPAC', R_PPAC(1),
       ' P_Lo', R_PLO(1),
       ' E Lo', R ELO(1),
       ' P Hi', R PHI(1),
       ' E_Hi',R_EHI(1))
       (A,F(7,0),A,F(3,0),A,F(10,3),A,F(5,0),A,F(6,0),A,F(10,3),A,F(6,0),A,F(10,3));
       @CALL U$PRTCL(C EVAP, U$M PRTTERM);
   END:
END;
IF (R IPAR(17) = 3) & (R PPAC(1) = 0) & (R SEL(1) > 3) THEN DO;
   IF (R ELO(1) > R IPAR(19)) & (R ELO(1) < R IPAR(20)) | ((R EHI(1) > 1850) & (R EHI(1)
< 2760)) THEN DO;
      PUT STRING(C EVAP) EDIT(
       'Ev',I_EVCT,
       ' St', R SEL(1),
       ' Time',TIME,
       ' P_Lo',R_PLO(1),
       ' E_Lo', R_ELO(1),
' P_Hi', R_PHI(1),
       ' E Hi', R EHI(1))
       (\texttt{A}, \texttt{F} (8, 0) \, , \texttt{A}, \texttt{F} (4, 0) \, , \texttt{A}, \texttt{F} (11, 3) \, , \texttt{A}, \texttt{F} (7, 0) \, , \texttt{A}, \texttt{F} (11, 3) \, , \texttt{A}, \texttt{F} (7, 0) \, , \texttt{A}, \texttt{F} (6, 0)) \, ;
       @CALL U$PRTCL(C EVAP, U$M PRTTERM);
   END;
END;
IF (R_IPAR(17) = 4) THEN DO;
   IF (I EVCT >= R IPAR(19)) & (I EVCT <= R IPAR(20)) THEN DO;
       I = R IPAR(18);
       PUT STRING(C EVAP) EDIT(
            I EVCT,
           TIMEMS,
           R PPAC(1),
           R SEL(1),
            R EL(I),
            R_PLO(1),
           R SEH(1),
            R EH(I),
            R PHI(1),
            R YHT(I),
            R_YHB(I),
            R YEHT(I),
            R YEHB(I))
            (\overline{F}(6), F(13), F(5), F(3), F(5), F(9,2), F(3), F(5), F(9,2), F(5), F(5), F(5), F(5));
       @CALL U$PRTCL(C EVAP, U$M PRTTERM);
   END;
END;
IF (R IPAR(17) = 5) THEN DO;
   IF (I\_EVCT >= R\_IPAR(19)) & (I\_EVCT <= R\_IPAR(20)) THEN DO;
       PUT STRING(C_EVAP) EDIT(
            I EVCT,
           TIMEUS,
           R PPAC(1),
            R SEL(1),
            R SYLT(1),
            R SYLB(1),
            R_CEL(1),
            R CYLT(1),
            R CYLB(1),
            R_PLO(1),
```

```
R SEH(1),
           R SYHT(1),
           R SYHB(1),
           R CEH(1),
           R CYHT(1),
           R CYHB(1),
           R PHI(1))
@CALL U$PRTCL(C EVAP, U$M PRTTERM);
END;
IF (R IPAR(17) = 6) & (I EVCT >= R IPAR(19)) & (I EVCT <= R IPAR(20)) THEN DO;
   DO^{-}I = 1 TO 32;
       PUT STRING(C_EVAP) EDIT(
           I,
           R EL(I),
           R YLT(I),
           R_YLB(I),
           R EH(I),
           R YHT(I),
           R_YHB(I))
           (\overline{F}(6), F(6), F(6), F(6), F(6), F(6), F(6));
       @CALL U$PRTCL(C EVAP, U$M PRTTERM);
   END;
END;
END EVENTLIST;
______
@RET(STS$VALUE);
    ENTRY CALLED DURING STARTUP OR BY COMMAND 'INITIALIZE ANALYSIS'
_______
$XANAL:ENTRY RETURNS(BIN FIXED(31));
@INCLUDE $MACRO($SECDEF);
/**** LOCATE DATA ELEMENTS *****/
$LOC(DE,DB,DATA,EVENT,W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SECAM =
P$_DB_DATA_EVENT;
$LOC(DE,DB,DATA,EL,W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P_EL = P$_DB_DATA_EL; $LOC(DE,DB,DATA,EH,W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P_EH = P$_DB_DATA_EH; $LOC(DE,DB,DATA,YLT,W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P_YLT =
                                          IF ^STS$SUCCESS THEN @RET(STS$VALUE); P_YLT =
$LOC(DE, DB, DATA, YLT, W);
P$ DB DATA YLT;
$LOC(DE, DB, DATA, YHT, W);
                                          IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YHT =
P$ DB DATA YHT;
$LOC (DE, DB, DATA, YLB, W);
                                          IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YLB =
P$_DB_DATA_YLB;
$LOC(DE, DB, DATA, YHB, W);
                                           IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YHB =
LY_DB_DATA_INB;

$LOC(DE, DB, DATA, BL, W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P_BL = P$_DB_DATA_BL;

$LOC(DE, DB, DATA, BH, W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P_BH = P$_DB_DATA_BH;

$LOC(DE, DB, DATA, PT, W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P_PT = P$_DB_DATA_PT;

$LOC(DE, DB, DATA, CEL, W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P_CEL =
P$ DB DATA CEL;
$LOC (DE, DB, DATA, CEH, W);
                                           IF ^STS$SUCCESS THEN @RET(STS$VALUE); P CEH =
P$ DB DATA CEH;
$LOC (DE, DB, DATA, CYLT, W);
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P CYLT =
P$_DB_DATA_CYLT;
```

```
$LOC(DE, DB, DATA, CYHT, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P CYHT =
P$ DB DATA CYHT;
$LOC(DE, DB, DATA, CYLB, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P CYLB =
P$ DB DATA CYLB;
$LOC(DE, DB, DATA, CYHB, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P CYHB =
P$ DB DATA CYHB;
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P CBL =
$LOC(DE, DB, DATA, CBL, W);
P$ DB DATA CBL;
$LOC(DE, DB, DATA, CBH, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P CBH =
P$ DB DATA CBH;
$LOC (DE, DB, DATA, CPT, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P CPT =
P$_DB_DATA_CPT;
$LOC (DE, DB, DATA, SEL, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SEL =
P$ DB DATA SEL;
$LOC(DE, DB, DATA, SEH, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SEH =
P$ DB DATA SEH;
$LOC(DE, DB, DATA, SYLT, W);
                              IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SYLT =
P$_DB_DATA_SYLT;
$LOC(DE, DB, DATA, SYHT, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SYHT =
P$ DB DATA_SYHT;
$LOC(DE, DB, DATA, SYLB, W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SYLB =
P$ DB DATA SYLB;
$LOC(DE, DB, DATA, SYHB, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SYHB =
P$ DB DATA SYHB;
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P_SBL =
$LOC(DE, DB, DATA, SBL, W);
P$ DB DATA SBL;
$LOC(DE, DB, DATA, SBH, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SBH =
P$ DB DATA SBH;
$LOC(DE, DB, DATA, SPT, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SPT =
P$ DB DATA SPT;
$LOC(DE, DB, DATA, ELM, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P ELM =
P$ DB DATA ELM;
$LOC (DE, DB, DATA, ELB, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P ELB =
P$ DB DATA ELB;
$LOC(DE, DB, DATA, EHM, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P EHM =
P$_DB_DATA_EHM;
$LOC(DE, DB, DATA, EHB, W);
                                       IF ^STS$SUCCESS THEN @RET(STS$VALUE); P EHB =
P$ DB DATA EHB;
$LOC(DE, DB, DATA, YLTM, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YLTM =
P$ DB_DATA_YLTM;
$LOC(DE, DB, DATA, YLTB, W);
                          IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YLTB =
P$ DB DATA YLTB;
$LOC(DE, DB, DATA, YLBM, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YLBM =
P$ DB DATA YLBM;
$LOC(DE,DB,DATA,YLBB,W);
                              IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YLBB =
P$ DB DATA YLBB;
$LOC(DE, DB, DATA, YHTM, W);
                              IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YHTM =
P$ DB DATA YHTM;
$LOC(DE, DB, DATA, YHTB, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YHTB =
P$ DB DATA YHTB;
$LOC(DE, DB, DATA, YHBM, W);
                              IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YHBM =
P$ DB DATA YHBM;
$LOC(DE, DB, DATA, YHBB, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YHBB =
P$_DB_DATA_YHBB;
$LOC(DE, DB, DATA, PKEV, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P PKEV =
P$_DB_DATA_PKEV;
$LOC(DE, DB, DATA, PMEV, W);
                              IF ^STS$SUCCESS THEN @RET(STS$VALUE); P PMEV =
P$_DB_DATA_PMEV;
$LOC (DE, DB, DATA, PLO, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P PLO =
P$ DB DATA_EHM;
$LOC(DE, DB, DATA, PHI, W);
                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE); P PHI =
P$ DB DATA_EHB;
$LOC(DE, DB, DATA, YELT, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YELT =
P$_DB_DATA_YELT;
$LOC(DE, DB, DATA, YEHT, W);
                             IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YEHT =
```

P\$ DB DATA YEHT;

```
$LOC(DE, DB, DATA, YELB, W);
                                IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YELB =
P$ DB DATA YELB;
$LOC (DE, DB, DATA, YEHB, W);
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P YEHB =
P$ DB DATA YEHB;
$LOC (DE, DB, DATA, EKEV, W);
                                IF ^STS$SUCCESS THEN @RET(STS$VALUE); P EKEV =
P$ DB DATA EKEV;
$LOC(DE,DB,DATA,EMEV,W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P EMEV =
P$ DB DATA EMEV;
$LOC(DE, DB, DATA, ELO, W);
                                          IF ^STS$SUCCESS THEN @RET(STS$VALUE); P ELO =
P$ DB DATA ELO;
                                          IF ^STS$SUCCESS THEN @RET(STS$VALUE); P EHI =
$LOC(DE, DB, DATA, EHI, W);
P$ DB DATA EHI;
$LOC(DE, DB, DATA, E EVR, W);
                                IF ^STS$SUCCESS THEN @RET(STS$VALUE); P E EVR =
P$ DB DATA E EVR;
$LOC(DE, DB, DATA, E MOM, W);
                                IF ^STS$SUCCESS THEN @RET(STS$VALUE); P E MOM =
P$ DB DATA E MOM;
$LOC (DE, DB, DATA, DTEA, W);
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P DTEA =
P$ DB DATA DTEA;
$LOC(DE, DB, DATA, E DAU, W);
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P E DAU =
P$ DB DATA E DAU;
$LOC(DE, DB, DATA, E EVRF, W);
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P E EVRF =
P$_DB_DATA_E_EVRF;
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P E FISS =
$LOC(DE, DB, DATA, E FISS, W);
P$ DB DATA E FISS;
$LOC(DE, DB, DATA, E PPAC, W);
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P E PPAC =
P$_DB_DATA_E_PPAC;
$LOC(DE, DB, DATA, NAIL, W);
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P NAIL =
P$ DB DATA NAIL;
$LOC (DE, DB, DATA, NAIH, W);
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P NAIH =
P$_DB_DATA_NAIH;
$LOC(DE,DB,DATA,RESLSW,W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P RESLSW =
P$ DB DATA RESLSW;
$LOC(DE, DB, DATA, RESMSW, W);
                                IF ^STS$SUCCESS THEN @RET(STS$VALUE); P RESMSW =
P$ DB DATA RESMSW;
$LOC(DE, DB, DATA, RWSLSW, W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P RWSLSW =
P$ DB DATA RWSLSW;
$LOC(DE,DB,DATA,RWSMSW,W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P RWSMSW =
P$ DB DATA RWSMSW;
$LOC(DE,DB,DATA,RE,W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P RE = P$ DB DATA RE; $LOC(DE,DB,DATA,RW,W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P RW = P$ DB DATA RW;
$LOC(DE, DB, DATA, IPAR, W);
                                IF ^STS$SUCCESS THEN @RET(STS$VALUE); P IPAR =
P$ DB DATA IPAR;
$LOC(DE, DB, DATA, SLOPE, W);
                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE); P SLOPE =
P$ DB DATA SLOPE;
$LOC(DE, DB, DATA, PPAC, W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P PPAC =
P$ DB DATA PPAC;
$LOC(DE, DB, DATA, ERROR, W); IF ^STS$SUCCESS THEN @RET(STS$VALUE); P ERROR =
P$ DB DATA ERROR;
/**** LOCATE CONDITIONS *****/
$LOC(COND, DB, $CONDITION, RE, W, WC);
                                              IF ^STS$SUCCESS THEN @RET(STS$VALUE);
IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(COND, DB, $CONDITION, RW, W, WC);
                                               IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(COND, DB, $CONDITION, PAUSE, W, WC);
                                                IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(COND, DB, $CONDITION, PPAC, W, WC);
                                                IF ^STS$SUCCESS THEN @RET(STS$VALUE);
IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(COND, DB, $CONDITION, E EVR, W, WC);
$LOC(COND, DB, $CONDITION, E_MOM, W, WC);
$LOC(COND, DB, $CONDITION, E_DAU, W, WC);
                                                IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(COND, DB, $CONDITION, E FISS, W, WC); IF ^STS$SUCCESS THEN @RET(STS$VALUE); $LOC(COND, DB, $CONDITION, E EVRF, W, WC); IF ^STS$SUCCESS THEN @RET(STS$VALUE); $LOC1(COND, DB, $CONDITION, PUNCH, 1, 8, W, WC); IF ^STS$SUCCESS THEN @RET(STS$VALUE);
/**** LOCATE SPECTRA ****/
$LOC1 (SPEC, DB, $SPECTRUM, S, 1, 352, W, L);
                                                 IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, SEC, W, L);
                                                  IF ^STS$SUCCESS THEN @RET(STS$VALUE);
```

```
$LOC(SPEC, DB, $SPECTRUM, TIME, W, L);
                                                                     IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                     IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, TIMEMIN, W, L);
$LOC(SPEC, DB, $SPECTRUM, RE RATE, W, L);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                     IF STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, RW RATE, W, L);
$LOC(SPEC, DB, $SPECTRUM, RUTH RATE, W, L);
                                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, TMP, W, L);
                                                                       IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC1(SPEC, DB, $SPECTRUM, SP, 1, 32, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, SPSUM, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, PPAC, W, L);
$LOC(SPEC, DB, $SPECTRUM, EVR_LOW, W, L);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, PLO, W, L);
$LOC1 (SPEC, DB, $SPECTRUM, PKEV, 1, 32, W, L);
                                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, PHI, W, L);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC1 (SPEC, DB, $SPECTRUM, PMEV, 1, 32, W, L);
                                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC1 (SPEC, DB, $SPECTRUM, EKEV, 1, 32, W, L);
$LOC(SPEC, DB, $SPECTRUM, SUML, W, L);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC1 (SPEC, DB, $SPECTRUM, EMEV, 1, 32, W, L);
                                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, SUMH, W, L);
                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC1 (SPEC, DB, $SPECTRUM, EPKEV, 1, 32, W, L);
                                                                   IF ^STS$SUCCESS THEN @RET(STS$VALUE);
IF ^STS$SUCCESS THEN @RET(STS$VALUE);
IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, SUMLP, W, L);
$LOC1 (SPEC, DB, $SPECTRUM, EPMEV, 1, 32, W, L);
$LOC(SPEC, DB, $SPECTRUM, SUMHP, W, L);
$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPEC, DB, $SPECTRUM, EAKEV, 1, 32, W, L);

$LOC1(SPECTRUM, 
$LOC(SPEC, DB, $SPECTRUM, SUMLA, W, L);
                                                                       IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC1 (SPEC, DB, $SPECTRUM, EAMEV, 1, 32, W, L); IF $ST$$SUCCESS THEN @RET(ST$$VALUE);
$LOC(SPEC,DB,$SFECTRUM,SUMHA,W,L); IF ^STS$SUCCESS THEN @RET(STS$VALUE); $LOC(SPEC,DB,$SPECTRUM,RUTHEAST,W,L); IF ^STS$SUCCESS THEN @RET(STS$VALUE); $LOC(SPEC,DB,$SPECTRUM,RUTHWEST,W,L); IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, SUMAPT, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, SUMPT, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, DT EA, W, L);
$LOC(SPEC, DB, $SPECTRUM, LOGDT EA, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, E EVAP, W, L);
                                                                     IF ^STS$SUCCESS THEN @RET(STS$VALUE);
IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, E MOTHER, W, L);
$LOC(SPEC, DB, $SPECTRUM, P MOTHER, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, P EVAP, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, DP EA, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, RATE EA, W, L);
$LOC(SPEC, DB, $SPECTRUM, E RATIO, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, DT AA, W, L);
$LOC(SPEC, DB, $SPECTRUM, E_EVAPALPHA, W, L);
                                                                      IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, E DAUGHTER, W, L);
                                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, DP AA, W, L);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                   IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, P EVAPALPHA, W, L);
$LOC(SPEC, DB, $SPECTRUM, P DAUGHTER, W, L);
                                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, DP EF, W, L);
$LOC(SPEC, DB, $SPECTRUM, E EVAPF, W, L);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, E FISSION, W, L);
                                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, P EVAPF, W, L);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, P FISSION, W, L);
                                                                                    IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, DT EF, W, L);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC(SPEC, DB, $SPECTRUM, RATE EF, W, L);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
                                                                        IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC1(SPEC, DB, $SPECTRUM, CT EVR, 1, 32, W, L);
$LOC1(SPEC, DB, $SPECTRUM, CT_MOM, 1, 32, W, L); IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC1(SPEC,DB,$SPECTRUM,CT_DAU,1,32,W,L); IF ^STS$SUCCESS THEN @RET(STS$VALUE); $LOC1(SPEC,DB,$SPECTRUM,CT_EVRF,1,32,W,L); IF ^STS$SUCCESS THEN @RET(STS$VALUE);
$LOC1 (SPEC, DB, $SPECTRUM, CT EVRF, 1, 32, W, L);
$LOC1(SPEC, DB, $SPECTRUM, CT FISS, 1, 32, W, L); IF ^STS$SUCCESS THEN @RET(STS$VALUE);
STS$VALUE=1;
@RET(STS$VALUE);
       PROCEDURE CALLED IN CASE OF AN ERROR
______
U CLEANUP: PROCEDURE;
END U CLEANUP;
```

/* 				 	 		
*/				 	 	======	
END /*	X\$ANAL;						
	END OF	PROCEDURE	X\$ANAL		 		
*/				 	 		

Appendix B: Patin parameters input file listing all of the HIVAP input parameters.

```
Ti-50 + Bi-209
                     Patin Parameters
 APROJ = 50 ZPROJ = 22 ATARG = 209 ZTARG = 83
 SHELL(GS) = 2 SHELF(SADDLE) = 2 PAIR = 4 MC = 0 MP = 0
 IBF = 1
 FISROT PARAMETERS = 0
 NOFISSION = 0 NONEUTRONS = 0 NOPROTONS = 0 NOALPHAS = 0 NOGAMMAS = 0
 DISC = 0 GAMMAS(IGAM) = 1
 NEUTRONS = 4 PROTONS = 1
 MASSES LOG UNIT = 9
 NUMB = 0 IOVER = 1 INERT = 0 INERF = 0 FINERT = 1
 LIMITS = 1
 PRINT = 5 \quad LOGUN = 0 \quad ICOR = 1
 NUMISO = 0 LBDM = 5
 IRAST = 0 WKB = 1 ITRANS = 0 JFJI = 1
______
ANG.MOM. LOSS NEUTRONS 1 PROTONS 1 ALPHAS 3
IF IGAM NOT ZERO
 EG1WU = 0.01 EG2WU = 10
 EG1MIN = 1 EG1MAX = 20
 EG2MIN = 1 EG2MAX = 4
 JFACT = 1
______
IF IGAM NOT ZERO
 CGIANT = 0.0 EGIANT = 80 WGIANT = 5 STRIPE = 0 IOPTG = 0
______
 LEVELPAR = 1.16 AF/AN = 2
 BARFAC = 1.0
 EDAMP = 18.0 BAR0 = 0.0
 SHELLO = 0.0 DELT= 11.0 QVALUE = 0.0
ONLY IF LDBM=(5)
 LDBM = 1 PREEX = 1 AX = 0 ENH = 0 BETA0 = 0 EDCOLL = 0 UCRIT = 5
ONLY IF LIMITS NOT ZERO
 CUT = 0.1E-6 FRACT2 = 0.1E-6
 ABSMIN = 0 PRCN = 0 SIGLOW = 0 DEL1 = 0 NOLEP = 0 NOLJI = 0 NOLJF = 0
______
IF INERT NOT ZERO (LOWER PART YRAST LINE)
 ENERGY SPIN ENERGY SPIN ENERGY
 E = 6 IEXC = 1 IFUS = 11 LIMBAR = 0
 JLOWER = 0 JUPPER = 0 NEWFIS = 0 TSTROT = 0 JFIS = 0 EN = 0
 V0 = 48 R0 = 1.12 D = 0.75 Q2 = 0 CRED = 1.0
 NOCURV = 0 NOPROX = 0 IOPT = 0 ITEST = 0
 SIGRO = 2.9 CUTOFF = 5.0 XTH = 0.7 APUSH = 18.0 FPUSH = 0.75
-1 0 0 0 0 0 0 0 0 0 0 0
```

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